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Patent application No. Demande de brevet no Patentanmeldung Nr.

03104280.7



Der Präsident des Europäischen Patentamts; Im Auftrag

For the President of the European Patent Offic

Le Président de l'Office européen des brevets p.o.

R C van Dijk

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Anmeldung Nr:

Application no.: 03104280.7

Demande no:

Anmeldetag:

Date of filing: 19.11.03

Date de dépôt:

Anmelder/Applicant(s)/Demandeur(s):

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Bezeichnung der Erfindung/Title of the invention/Titre de l'invention: (Falls die Bezeichnung der Erfindung nicht angegeben ist, siehe Beschreibung. If no title is shown please refer to the description. Si aucun titre n'est indiqué se referer à la description.)

Seedyl sequence for making plants having changed growth characteristics

In Anspruch genommene Prioriät(en) / Priority(ies) claimed /Priorité(s) revendiquée(s)
Staat/Tag/Aktenzeichen/State/Date/File no./Pays/Date/Numéro de dépôt:

Internationale Patentklassifikation/International Patent Classification/Classification internationale des brevets:

C07K14/00

Am Anmeldetag benannte Vertragstaaten/Contracting states designated at date of filing/Etats contractants désignées lors du dépôt:

AT BE BG CH CY CZ DE DK EE ES FI FR GB GR HU IE IT LU MC NL PT RO SE SI SK TR LI

# Seedy1 sequence for making plants having changed growth characteristics

The present invention concerns a method for modifying growth characteristics of a plant. More specifically, the present invention concerns a method for modifying growth characteristics by modified expression of a seedy1 nucleic acid and/or by modified levels and/or activity of a seedy1 protein in a plant. The present invention also concerns plants having modified expression of a seedy1 nucleic acid and/or modified levels and/or activity of a seedy1 protein, which plants have modified growth characteristics relative to corresponding wild type plants.

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The ever-increasing world population and the dwindling supply of arable land available for agriculture fuel research towards improving the efficiency of agriculture. Conventional means for crop and horticultural improvements utilise selective breeding techniques to identify plants having desirable characteristics. However, such selective breeding techniques have several drawbacks, namely that these techniques are typically labour intensive and result in plants that often contain heterogeneous genetic components that may not always result in the desirable trait being passed on from parent plants. Advances in molecular biology have allowed mankind to modify the germplasm of animals and plants. Genetic engineering of plants entails the isolation and manipulation of genetic material (typically in the form of DNA or RNA) and the subsequent introduction of that genetic material into a plant. Such technology has the capacity to deliver crops or plants having various improved economic, agronomic or horticultural traits. A trait of particular economic interest is yield. Yield is normally defined as the measurable produce of economic value from a crop. This may be defined in terms of quantity and/or quality. Crop yield may not only be increased by combating one or more of stresses to which a crop or plant is typically subjected, but may also be increased by modifying the inherent growth characteristics of a plant. Yield is directly dependent on several growth characteristics, for example, the growth rate, the biomass production, plant architecture, number and size of the organs, (for example, the number of branches, tillers, shoots, flowers), seed production and more.

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The ability to influence one or more of the abovementioned growth characteristics, would have many applications in areas such as crop enhancement, plant breeding, production of ornamental plants, aboriculture, horticulture, forestry, production of algae or plants (for example for use as bioreactors, for the production of substances such as pharmaceuticals, antibodies, or vaccines, or for the bioconversion of organic waste or for use as fuel in the case of high-yielding algae and plants).

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It has now been found that modification in a plant of the expression of a seedy1 nucleic acid and/or modification of the level and/or activity levels in a plant of a seedy1 protein, gives rise to plants having modified growth characteristics. It has been shown that introduction of a seedy1 nucleic acid effects an increase in above ground biomass, increased total seed weight, increased number of seeds and increased number of tillers. It has been found that the hitherto unknown protein seedy1, is a protein having a coiled coil domain and three conserved motifs as presented in SEQ ID NO's 15, 16 and 17.

Therefore, the present invention provides a method for modifying growth characteristics of a plant, comprising modifying in a plant expression of a nucleic acid encoding a seedy1 protein and/or modifying in a plant level and/or activity of a seedy1 protein, wherein said seedy1 protein comprises in the following order from N-terminus to C-terminus at least two motifs, which motifs are:

- (i) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 15; and/or
- (ii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 16, and/or
- (iii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 17 and which is a coiled coil motif; and/or
- (iv) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 18.

and wherein said growth characteristics are modified relative to the corresponding wildtype plants.

Modifying expression of a seedy1 nucleic acid and/or modifying of the activity and/or levels of a seedy1 protein encompasses modifying expression of a gene and/or modifying acti vity and/or levels of a gene product, namely a polypeptide, in specific cells or tissues. The term "modifying" as used herein means increasing, decreasing or changing in time or place. The modified expression, activity and/or levels of a seedy1 gene or protein are modified compared to expression, activity and/or levels of a seedy1 gene or protein in corresponding wild-type plants. The modified gene expression may result from modified expression levels of an endogenous seedy1 gene and/or may result from modified expression levels of a seedy1 gene previously introduced into a plant. Similarly, modified levels and/or activity of a seedy1 protein may be due to modified expression of an endogenous seedy1 nucleic acid/gene and/or due to modifyed expression of a seedy1 nucleic acid/gene previously introduced into a plant. Modified expression of a gene/nucleic acid and/or modifying activity and/or levels of a gene product/protein may be effected, for example, by chemical means and/or recombinant means.

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Therefore there is provided by the present invention, a method for modifying growth characteristics of a plant, comprising modifying seedy1 gene expression and/or modifying seedy1 protein levels and/or seedy1 protein activity, which modification may be effected by recombinant means and/or by chemical means.

Advantageously, modified expression of a seedy1 nucleic acid and/or modified activity and/or levels of a seedy1 protein may be effected by chemical means, i.e. by exogenous application of one or more compounds or elements capable of modifying expression of a seedy1 nucleic acid and/or capable of modifying activity and/or levels of the seedy1 protein. The term "exogenous application" as defined herein is taken to mean the contacting or administering of a suitable compound or element to a plant (as defined herein below, the term "plant" includes plant cell, tissue, organ or to the whole organism). The compound or element may be exogenously applied to a plant in a form suitable for plant uptake (such as through application to the soil for uptake via the roots, or in the case of some plants by applying directly to the leaves, for example by spraying). The exogenous application may take place on wild-type plants or on transgenic plants that have previously been transformed with a seedy1 nucleic acid/gene or another transgene.

- 20 Suitable compounds or elements include seedy1 proteins or seedy1 nucleic acids. Alternatively, exogenous application of compounds or elements capable of modifying levels of factors that directly or indirectly activate or inactivate a seedy1 protein will also be suitable in practising the invention. Also included are antibodies that can recognise or mimic the function of seedy1 proteins. Such antibodies may comprise "plantibodies", single chain antibodies, IgG 25 antibodies and heavy chain camel antibodies, as well as fragments thereof. Additionally or alternatively, the resultant effect may also be achieved by the exogenous application of an interacting protein or activator or an inhibitor of the seedy1 gene/gene product. Additionally or alternatively, the compound or element may be a mutagenic substance, such as a chemical selected from any one or more of: N-nitroso-N-ethylurea, ethylene imine, ethyl methanesulphonate and diethyl sulphate. Mutagenesis may also be achieved by exposure to 30 ionising radiation, such as X-rays or gamma-rays or ultraviolet light. Methods for introducing mutations and for testing the effect of mutations (such as by monitoring gene expression and/or protein activity) are well known in the art.
- Therefore, according to one aspect of the present invention, there is provided a method for modifying growth characteristics of a plant, comprising exogenous application of one or more

compounds or elements capable of modifying expression of a nucleic acid encoding a seedy1 protein and/or capable of modifying activity and/or levels of a seedy1 protein.

Additionally or alternatively, and according to a preferred embodiment of the present invention, modification of expression of a seedy1 nucleic acid and/or modification of activity and/or levels of a seedy1 protein may be effected by recombinant means. Such recombinant means may comprise a direct and/or indirect approach for modification of expression of a nucleic acid encoding a seedy 1 and/or for modification of the activity and/or levels of a seedy1 protein.

The nucleic acid encoding a seedy one protein or the seedy1 protein as mentioned above may be wild type, i.e. a native or endogenous nucleic acid or protein. Alternatively, it may be a nucleic acid derived from the same or another species, which nucleic acid is introduced as a transgene, for example by transformation. This transgene may be substantially changed from its native form in composition and/or genomic environment through deliberate human manipulation.

An indirect recombinant approach may comprise for example introducing, into a plant, a nucleic acid capable of modifying expression of the gene in question (a seedy1 gene) and/or capable of modifying activity and/or levels of the protein in question (a seedy1 protein). Examples of such nucleic acids to be introduced into a plant are nucleic acids encoding transcription factors or activators or inhibitors that bind to the promoter of a seedy1 gene or that interact with a seedy1 protein. Methods to test these types of interactions and methods for isolating nucleic acids encoding such interactors include yeast one-hybrid or a yeast two-hybrid screens wherein the seedy1 gene/protein is used as a bait.

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Also encompassed by an indirect approach for modifying expression of a seedy1 gene and/or activity and/or levels of a seedy1 protein, is the provision of, or the inhibition or stimulation of regulatory sequences that drive expression of the native seedy1 gene or of the seedy1 transgene. Such regulatory sequences may be introduced into a plant. For example, the nucleic acid introduced into the plant is a promoter, capable of driving the expression of an endogenous seedy1 gene.

A further indirect approach for modifying expression of a seedy1 gene and/or for modifying activity and/or levels of a seedy1 protein in a plant, encompasses modified levels in a plant of a factor able to interact with seedy1. Such factors may include ligands of seedy1. Therefore, the present invention provides a method for modifying growth characteristics of a plant, comprising modifying expression of a gene coding for a protein which is a natural ligand of a

seedy1. Furthermore, the present invention also provides a method for modifying grow characteristics of a plant, comprising modifying expression of a gene coding for a protein which is a natural target/substrate of a seedy1.

A direct and more preferred approach for modifying growth characteristics of a plar comprises introducing into a plant a seedy1 nucleic acid, or a portion thereof or sequence capable of hybridising therewith, which nucleic acid preferably encodes a seedy1 protein or homologue, derivative or active fragment thereof. The nucleic acid may be introduced into plant by, for example, transformation.

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Accordingly, the present invention provides a method for modifying growth characteristics of plant, comprising introducing into a plant a nucleic acid capable of modifying expression of nucleic acid encoding a seedy1 protein and/or capable of modifying activity and/or levels of seedy1 protein. Further preferably such nucleic acid is a seedy1 nucleic acid.

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As mentioned above the nucleic acid to be used in the methods of the present invention ca be wild type (native or endogenous). Alternatively, the nucleic acid may be derived from another species, which gene is introduced into the plant as a transgene, for example t transformation. The nucleic acid may thus be derived (either directly or indirectly subsequently modified)) from any source provided that the nucleic acid, when expressed in plant, leads to modified expression of a seedy1 nucleic acid/gene or modified activity and/k levels of a seedy1 protein. The nucleic acid may be isolated from a microbial source, such a bacteria, yeast or fungi, or from a plant, algae, insect, or animal (including human) source Preferably, the seedy1 nucleic acid is isolated from a plant. This nucleic acid may b substantially changed from its native form in composition and/or genomic environment through deliberate human manipulation. The nucleic acid may be isolated from a dicotyledonou species, preferably from the family Solanaceae, further preferably from Nicotiana. Mor preferably, the nucleic acid is as represented by SEQ ID NO: 1 or a portion thereof or a nuclei acid capable of hybridising therewith or is a nucleic acid encoding an amino acid represente by SEQ ID NO: 2 or a homologue derivative or active fragment thereof, such as a homologu having at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80% 85%, 90%, 95% 98%, 99% sequence identity with SEQ ID NO 2.

Advantageously, the methods according to the invention may also be practised using variar nucleic acids and variant amino acids of SEQ ID NO 1 or 2 respectively, such as the variant further defined hereinafter.

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The term "seedy1 gene" or "seedy1 nucleic acid" or "nucleic acid encoding a seedy1 protein" are used herein interchangeably.

Taken in a broad context, the term "seedy" protein/nucleic acid also encompasses variant nucleic acids and variant amino acids suitable for practicing the methods according to the invention. Preferably, variant nucleic acids and variant amino acids suitable for practicing the methods according to the invention include those falling within the definition of a "seedy1" meaning a encoding a protein ore being a protein comprising in the following order from N-terminus to C-terminus at least two motifs, which motifs are:

- (i) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 15; and/or
- (ii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 16, and/or
- (iii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 17 and which is a coiled coil motif; and/or
- (iv) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 18.

For determining the presence of these conserved motifs sequence can be aligned with the software such as for example Align X or clustal X, for indication of the conserved residues (see for example Figure 3). Software packages like MEME version 3.0 can also be used to determine motifs in sequences. This software is available from UCSD, SDSC and NBCR at http://meme.sdsc.edu/meme/. For the identification of a coiled coil domain, the software Coils 2.0 can be used. This software is available at <a href="http://www.ch.embnet.org/software/COILS form.html">http://www.ch.embnet.org/software/COILS form.html</a>. For the presentation of the motifs in SEQ Id NO 15, 16, 17 and 18, X represents any amino acid.

According to a particular embodiment a seedy 1 protein has at least two, preferably three, most preferably four of the above defined domains.

- 30 According to a particular embodiment of the invention, a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 15 is the core sequence WXNAXXD as represented in SEQ ID NO 15, at least the or part of sequence (P/X)X((V/L/H)(Q/H)(V/I)W(N/X)NA(A/P)(F/C)D wherein
  - (P/X) preferably is P or is A or T or Q or another amino acid
- 35 (V/L/H) preferable is V or L or H
  - (Q/H) is either Q or H
  - (V/I) is either V or is T or S or another amino acid

(A/P) is preferable A or is P

(F/C) is preferably F or is C

Alternatively or additionally, according to one embodiment a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 16, is the core sequence KENXXP as represented in SEQ ID NO 16.

Alternatively or additionally, according to one embodiment a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 17 and which is a coiled coil motif is the core sequence EX<sub>1-8</sub>EXXRLXXXLXXLR as represented in SEQ ID NO 17, or is at least part of the sequence

(I/V/A)(D/E)XE(I/M)XX(I/V)(E/Q)XE(I/X)XRL(S/X)(S/X)(R/K)LXXLR(L/V/T/I)X(K/Q), wherein (I/V/A) preferable is I or V or is A

(D/E) is either D or E

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15 (I/M) preferably is I or is M

(I/V) preferably is I or is V

(E/Q) preferably is E or is Q

(I/X) preferably is I or is M or is V or any other amino acid

(S/X) preferably S or is T or any other amino acid

20 (S/X) preferably is S or isT or L or I or A

(R/K) preferably is R or is K

(L/V/T/I) preferable is L or T or V or I

(K/Q) preferably is K or Q

Alternatively or additionally, according to one embodiment a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 18, is the core sequence LPXIX<sub>1-10</sub>RDSGXXKRX<sub>1-6</sub>K as represented in SEQ ID NO 18, or is at least part of the sequence

.30 .. /X)(G/A)K, wherein .......

(R/K) is either R or K

(R/X) is preferably R or is S or K

(T/I) is preferably T or I

(M/X) is preferably M or L or A or V

35 (P/R) is either P or R

(D/X) is preferably D or is G or T or N

(E/G) is preferably E or is G

(S/T) is preferably S or is T

(P/L) is preferably P or is L

(C/X) is preferably C or is P or A

(A/X) is preferably A or is V or I

5 (A/I) is preferably A or is I

(D/E) is either D or E

(L/R) is preferably L or is R

(V/X) is preferably V or is Q or N or I

(G/A) is preferably G or is A

According to a further embodiment the nucleic acid has a motif which has at least 80% sequence identity to the sequence RDSGXXKRX<sub>1-6</sub>K.

Examples of such seedy1 proteins are monocots seedy1 proteins such as represented by SEQ ID NO 4 (rice), SEQ ID NO 8 (sugar cane) and SEQ ID NO 10 (maize) or from dicots SEQ ID NO 2 (tobacco), SEQ ID NO 6 (medicago) or SEQ ID NO 12 (Arabidopsis). The proteins as presented in SEQ ID NO 8 (sugar cane) and SEQ ID NO 10 (Maize) are only partial, but the corresponding full length sequences of the proteins and encoding cDNA, are well within the reach of a person skilled in the art who is now able to perform colony hybridization of a cDNA library, or PCR based on the use of specific primers combined with degenerated primers.

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Suitable variant nucleic acid and amino acid sequences useful in practising the method according to the invention, include:

- (i) Functional portions of a seedy1 nucleic acid/gene;
- (ii) Sequences capable of hybridising with a seedy1 nucleic acid/gene;
- (iii) Alternative splice variants of a seedy1 nucleic acid/gene;
- (iv) Allelic variants of a seedy1 nucleic acid/gene;
- (v) Homologues, derivatives and active fragments of a seedy1 protein;

The term seedy1 nucleic acid/gene, as defined herein, also encompasses a complement of SEQ ID NO 1 and also to corresponding RNA, DNA, cDNA or genomic DNA. The seedy1 may be synthesized in whole or in part, it may be double-strand nucleic acid or single-stranded nucleic acid. Also this term encompasses a variant of the gene due to the degeneracy of the genetic code and variants that are interrupted by one or more intervening sequences.

An example of a variant seedy1 nucleic acid is a functional portion of a seedy1 nucleic acid. The methods according to the invention may advantageously be practised using functional portions of a seedy1. A functional portion refers to a piece of DNA derived or prepared from

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an original (larger) DNA molecule, which DNA portion, when introduced and expressed in plant, gives plants having changed development. The portion may comprise many genes, will or without additional control elements or may contain spacer sequences. The portion may be made by making one or more deletions and/or truncations to the nucleic acid. Techniques for introducing truncations and deletions into a nucleic acid are well known in the art. Portion suitable for use in the methods according to the invention may readily be determined be following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the portion to be tested for functionality.

- An example of a further variant seedy1 nucleic acid is a sequence that is capable of hybridising to a seedy1 nucleic acid, for example to SEQ ID NO 1, 3, 5, 7, 9 or 11 Advantageously, the methods according to the present invention may also be practised using sequences capable of hybridising to a coiled coil protein, particularly a seedy1 protein as represented by any one of SEQ ID NO: 2, 4, 6, 8, 10 or 12, which hybridising sequences are preferably those falling within the definition of a "seedy1" as set out herein before. Hybridising sequences suitable for use in the methods according to the invention may readily be determined for example by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the hybridising sequence.
- The term "hybridisation" as defined herein is a process wherein substantially homologous 20 complementary nucleotide sequences anneal to each other. The hybridisation process can occur entirely in solution, i.e. both complementary nucleic acids are in solution. Tools in molecular biology relying on such a process include the polymerase chain reaction (PCR; and all methods based thereon), subtractive hybridisation, random primer extension, nuclease S1 25 mapping, primer extension, reverse transcription, cDNA synthesis, differential display of RNAs, and DNA sequence determination. The hybridisation process can also occur with one of the complementary nucleic acids immobilised to a matrix such as magnetic beads, Sepharose beads or any other resin. Tools in molecular biology relying on such a process include the isolation of poly (A+) mRNA. The hybridisation process can furthermore occur with one of the complementary nucleic acids immobilised to a solid support such as a nitro-cellulose or nylon-30 membrane or immobilised by e.g. photolithography to e.g. a siliceous glass support (the latter known as nucleic acid arrays or microarrays or as nucleic acid chips). Tools in molecular biology relying on such a process include RNA and DNA gel blot analysis, colony hybridisation, plaque hybridisation, in situ hybridisation and microarray hybridisation. In order to allow hybridisation to occur, the nucleic acid molecules are generally thermally or chemically 35 denatured to melt a double strand into two single strands and/or to remove hairpins or other secondary structures from single stranded nucleic acids. The stringency of hybridisation is

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influenced by conditions such as temperature, salt concentration and hybridisation buffer composition. High stringency conditions for hybridisation include high temperature and/or low salt concentration (salts include NaCl and Na<sub>3</sub>-citrate) and/or the inclusion of formamide in the hybridisation buffer and/or lowering the concentration of compounds such as SDS (detergent) in the hybridisation buffer and/or exclusion of compounds such as dextran sulphate or polyethylene glycol (promoting molecular crowding) from the hybridisation Conventional hybridisation conditions are described in, for example, Sambrook (2001) Molecular Cloning: a laboratory manual, 3rd Edition Cold Spring Harbor Laboratory Press, CSH, New York, but the skilled craftsman will appreciate that numerous different hybridisation conditions can be designed in function of the known or the expected homology and/or length of the nucleic acid. Sufficiently low stringency hybridisation conditions are particularly preferred (at least in the first instance) to isolate nucleic acids heterologous to the DNA sequences of the invention defined supra. An example of low stringency conditions is 4-6x SSC / 0.1-0.5% w/v SDS at 37-45°C for 2-3 hours. Depending on the source and concentration of the nucleic acid involved in the hybridisation, alternative conditions of stringency may be employed, such as medium stringency conditions. Examples of medium stringency conditions include 1-4x SSC / 0.25% w/v SDS at ≥ 45°C for 2-3 hours. An example of high stringency conditions includes 0.1-1x SSC / 0.1% w/v SDS at 60°C for 1-3 hours. The skilled man will be aware of various parameters which may be altered during hybridisation and washing and which will either maintain or change the stringency conditions. The stringency conditions may start low and be progressively increased until there is provided a hybridising seedy1 nucleic acid, as defined hereinabove. Elements contributing to heterology include allelism, degeneration of the genetic code and differences in preferred codon usage.

Another example of a variant seedy1 is an alternative splice variant of a seedy1. The methods according to the present invention may also be practised using an alternative splice variant of a seedy1 nucleic acid. The term "alternative splice variant" as used herein encompasses variants of a nucleic acid in which selected introns and/or exons have been excised, replaced or added. Such splice variants may be found in nature or can be manmade using techniques well known in the art. Preferably, the splice variant is a splice variant of the sequence represented by any of SEQ ID NO 1, 3, 5, 7, 9 or 11. Splice variants suitable for use in the methods according to the invention may readily be determined for example by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the splice variants.

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Another example of a variant seedy1 is an allelic variant. Advantageously, the methods according to the present invention may also be practised using allelic variants of a seedy1

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nucleic acid, preferably an allelic variant of a sequence represented by any of SEQ ID NO 1, 3, 5, 7, 9 or 11. Allelic variants exist in nature and encompassed within the methods of the present invention is the use of these isolated natural alleles in the methods according to the invention. Allelic variants encompass Single Nucleotide Polymorphisms (SNPs), as well as Small Insertion/Deletion Polymorphisms (INDELs). The size of INDELs is usually less than 100 bp). SNPs and INDELs form the largest set of sequence variants in naturally occurring polymorphic strains of most organisms. Allelic variants suitable for use in the methods according to the invention may readily be determined for example by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the allelic variant.

Accordingly, the present invention provides a method for modifying growth characteristics of a plant, wherein the seedy1 nucleic acid is a splice variant of a seedy1 nucleic acid or wherein said seedy1 protein is encoded by a splice variant or wherein the seedy1 nucleic acid is an allelic variant of a seedy1 nucleic acid or wherein said seedy1 protein is encoded by an allelic variant.

Examples of variant seedy1 amino acids also in clude homologues, derivatives and active fragments of a seedy1 protein. Advantageously, the methods according to the present invention may also be practised using homologues, derivatives or active fragments of a coiled coil, preferably using homologues, derivatives or active fragments of a seedy1 protein, preferably a seedy1 protein as represented by any one of SEQ ID NO 2, 4, 6, 8, 10 or 12.

"Homologues" of a seedy1 protein encompass peptides, oligopeptides, polypeptides, proteins and enzymes having amino acid substitutions, deletions and/or insertions relative to the unchanged protein in question and having similar biological and functional activity as the unchanged protein from which they are derived. To produce such homologues, amino acids of the protein may be replaced by other amino acids having similar properties (such as similar hydrophobicity, hydrophilicity, antigenicity, propensity to form or break  $\alpha$ -helical structures or  $\beta$ -sheet structures). Conservative substitution tables are well known in the art (see for example Creighton (1984) Proteins. W.H. Freeman and Company).

The homologues useful in the methods according to the invention have a percentage identity to any one of SEQ ID NO 2, 4, 6, 8, 10 or 12 equal to value lying between 20% and 99.99%.

The homologues useful in the method according to the invention have at least 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, or 50% sequence identity or

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similarity (functional identity) to the unchanged protein, alternatively at least 60% sequence identity or similarity to an unchanged protein, alternatively at least 70% sequence identity or similarity to an unchanged protein. Typically, the homologues have at least 75% or 80% sequence identity or similarity to an unchanged protein, preferably at least 85%, 86%, 87%, 88%, 98% sequence identity or similarity, further preferably at least 90%, 91%, 92%, 93%, 94% sequence identity or similarity to an unchanged protein, most preferably at least 95%, 96%, 97%, 98% or 99% sequence identity or similarity to an unchanged protein.

The percentage of identity can be calculated by using an alignment program well known in the art. For example, the percentage of identity can be calculated using the program GAP, or needle (EMBOSS package) or stretcher (EMBOSS package) or the program align X, as a module of the vector NTI suite 5.5 software package, using the standard parameters (for example GAP penalty 5, GAP opening penalty 15, GAP extension penalty 6.6).

The homologues useful in the methods according to the invention are preferably coiled coil proteins, further preferably seedy1 proteins as defined herein above. Homologues suitable for use in the methods according to the invention may readily be determined for example by following the methods described in the Examples section by simply substituting the sequence used in the actual Example with the homologous sequence.

Methods for the search and identification of seedy1 homologues or DNA sequences encoding a seedy1 homologue, would be well within the realm of persons skilled in the art. Such methods, involve screening sequence databases with the sequences as provided by the present invention in SEQ ID NO 1 and 2 or 3 to 10, preferably a computer readable format of the nucleic acids of the present invention. This sequence information is available for example to Genbank public databases, that include but are not limited in (http://www.ncbi.nlm.nih.gov/web/Genbank), the European Molecular Biology Laboratory Nucleic acid Database (EMBL) (http://w.ebi.ac.uk/ebi-docs/embl-db.html) or versions thereof or the MIPS database (http://mips.gsf.de/). Different search algorithms and software for the alignment and comparison of sequences are well known in the art. Such software includes software include GAP, BESTFIT, BLAST, FASTA and TFASTA. GAP uses the algorithm of Needleman and Wunsch (J. Mol. Biol. 48: 443-453, 1970) to find the alignment of two complete sequences that maximises the number of matches and minimises the number of gaps. The BLAST algorithm calculates percentage sequence identity and performs a statistical analysis of the similarity between the two sequences. The suite of programs referred to as BLAST programs has 5 different implementations: three designed for nucleotide sequence queries (BLASTN, BLASTX, and TBLASTX) and two designed for protein sequence queries (BLASTP and TBLASTN) (Coulson, Trends in Biotechnology: 76-80, 1994; Birren et al.,

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GenomeAnalysis, 1: 543, 1997). The software for performing BLAST analysis is public available through the National Centre for Biotechnology Information.

Homologues of SEQ ID NO 2 can be found in many different organisms. The closes homologues are found in the plant kingdom. For example, seedy1 nucleic acids were isolate from tobacco (SEQ ID NO 1 and 2), from rice (SEQ ID NO 3) encoding a rice seedy homologue (SEQ ID NO 4), from medicago (SEQ ID NO 5 and 6), from sugar cane (SEQ IE NO 7 and 8), from maize (SEQ ID NO 9 and 10) and from Arabidopsis (SEQ ID NO 11 and 12). Also EST's from other organisms were found and deposited in Genbank, for example ES1 from Vitis vinifera (accession number CA816066), from Pinus taeda (accession number BM903108), from Saccharus sp. (So variant) (accession numbers CA228193 and CA256020) from Citrus sinsensis (accession number CF833583), Plumbago zeylanica (accession number CB817788), from Zea mays (accession number CF637447, AW282224, CD058812, AY108162, CD059048, CF041861, AW067243), from Triticum aestivum (CA727065, BJ264506, BJ259034), from Hordeum vulgare (accession number BU997034, CA727065, CA031127, BQ762011), from Bn (CD817460) from Ga (BG446106, BM360339), from Eschscholzia califronica (CD478368), from Pt (BU821376), and from beta vulgaris (BQ594009).

As more genomes are being sequenced, it is expected that many more seedy1 homologues shall be identifiable.

The above-mentioned analyses for comparing sequences, for calculation of sequence identity and for the search for homologues, is preferentially done with the full-length sequence or with a conserved region of the sequence. Therefore, these analyses can be based on a comparison of certain regions such as conserved domains, motifs or boxes.

The identification of such domains or motifs, would also be well within the realm of a person skilled in the art and involves for example, a computer readable format of the nucleic acids of the present invention, the use of alignment software programs and the use of publicly available information on protein domains, conserved motifs and boxes. This protein domain information

is .. . .. . .30. . · · · available the **PRODOM** (http://www.biochem.ucl.ac.uk/bsm/dbbrowser/jj/prodomsrchjj.html), PIR (http://pir.georgetown.edu/) or pFAM (http://pfam.wustl.edu/) database. Sequence analysis programs designed for motif searching can be used for identification of fragments, regions and conserved domains as mentioned above. Preferred computer programs would include but are not limited to MEME, SIGNALSCAN, and GENESCAN. A MEME algorithm (Version 3.0) can be found in the GCG package; or on the Internet site http://www.sdsc.edu/MEME/meme. SIGNALSCAN version 4.0 information available is on the Internet site

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http://biosci.cbs.umn.edu/software/sigscan.html. GENESCAN can be found on the Internet site http://gnomic.stanford.edu/GENESCANW.html.

More particularly preferred seedy1 homologues have the conserved domains as depicted in SEQ ID NO 15, 16 and 17, or motifs that are 80% identical to these motifs. Also preferred seedy1 homologues have a coiled coil domain, preferably located in the N-terminal half of the protein, more preferably between amino acid position 25 to 250, more preferably between position 50 and 150. A coiled coil domain is determining the folding of the protein.

Two special forms of homology, orthologous and paralogous, are evolutionary concepts used to describe ancestral relationships of genes. The term "paralogous" relates to geneduplications within the genome of a species. A "paralogue" is therefore duplication. The term "orthologous" relates to homologous genes in different organisms due to ancestral relationship and the formation of different species. An orthologue is thus a "speciation". The term "homologues" as used herein also encompasses paralogues and orthologues and are useful proteins in the methods according to the invention.

Another variant of seedy1 useful in the methods of the present invention is a derivative of seedy1. The term "derivatives" refers to peptides, oligopeptides, polypeptides, proteins and enzymes which may comprise substitutions, deletions or additions of naturally and non-naturally occurring amino acid residues compared to the amino acid sequence of a naturally-occurring form of the protein, for example, as presented in SEQ ID NO: 2. "Derivatives" of a seedy1 protein encompass peptides, oligopeptides, polypeptides, proteins and enzymes which may comprise naturally occurring changed, glycosylated, acylated or non-naturally occurring amino acid residues compared to the amino acid sequence of a naturally-occurring form of the polypeptide. A derivative may also comprise one or more non-amino acid substituents compared to the amino acid sequence from which it is derived, for example a reporter molecule or other ligand, covalently or non-covalently bound to the amino acid sequence such as, for example, a reporter molecule which is bound to facilitate its detection, and non-naturally occurring amino acid residues relative to the amino acid sequence of a naturally-occurring protein.

"Substitutional variants" of a protein are those in which at least one residue in an amino acid sequence has been removed and a different residue inserted in its place. Amino acid substitutions are typically of single residues, but may be clustered depending upon functional constraints placed upon the polypeptide; insertions will usually be of the order of about 1 to 10

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amino acid residues, and deletions will range from about 1 to 20 residues. Preferably, amino acid substitutions comprise conservative amino acid substitutions.

"Insertional variants" of a protein are those in which one or more amino acid residues are introduced into a predetermined site in a protein. Insertions can comprise amino-terminal and/or carboxy-terminal fusions as well as intra-sequence insertions of single or multiple amino acids. Generally, insertions within the amino acid sequence will be smaller than amino- or carboxy-terminal fusions, of the order of about 1 to 10 residues. Examples of amino- or carboxy-terminal fusion proteins or peptides include the binding domain or activation domain of a transcriptional activator as used in the yeast two-hybrid system, phage coat proteins, (histidine)6-tag, glutathione S-transferase-tag, protein A, maltose-binding protein, dihydrofolate reductase, Tag·100 epitope, c-myc epitope, FLAG®-epitope, lacZ, CMP (calmodulin-binding peptide), HA epitope, protein C epitope and VSV epitope.

"Deletion variants" of a protein are characterised by the removal of one or more amino acids from the protein. Amino acid variants of a protein may readily be made using peptide synthetic techniques well known in the art, such as solid phase peptide synthesis and the like, or by recombinant DNA manipulations. Methods for the manipulation of DNA sequences to produce substitution, insertion or deletion variants of a protein are well known in the art. For example, techniques for making substitution mutations at predetermined sites in DNA are well known to those skilled in the art and include M13 mutagenesis, T7-Gen in vitro mutagenesis (USB, Cleveland, OH), QuickChange Site Directed mutagenesis (Stratagene, San Diego, CA), PCR-mediated site-directed mutagenesis or other site-directed mutagenesis protocols.

Another variant of seedy1 useful in the methods of the present invention is an active fragment of seedy1. "Active fragments" of a seedy1 protein encompasses contiguous amino acid residues of a seedy1 protein, which residues retain similar biological and/or functional activity to the naturally occurring protein. For example, useful fragments comprise at least 10 contiguous amino acid residues of a seedy1 protein. Other preferred fragments are fragments of the seedy1 protein starting at the second or third or further internal methionin residues. These fragments originate from protein translation, starting at internal ATG codons.

In the present invention a new category of proteins and encoding genes is disclosed. Surprisingly, altering the availability of such sequences is a plant changes the growth characteristics of the plant. More particularly the seed quantity and quality is improved in these plants. Therefore, these sequences useful for the methods of the present invention are named "seedy" sequences.

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As the type of proteins defined herein as seedy1 proteins was not known, the present invention provides for an isolated nucleic acid encoding at least part of a seedy1 protein, wherein said seedy1 protein comprises in the following order from N-terminus to C-terminus at least two motifs, which motifs are:

- (i) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 15; and/or
- (ii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 16, and/or
- (iii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 17 and which is a coiled coil motif; and/or
- (iv) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 18.
- In the present invention, a new type of gene has been isolated from tobacco which gene encodes a new type of protein. Surprisingly it has been found that this gene and/or protein is capable of modifying growth characteristics of a plant. Therefore, the present invention provides an isolated nucleic acid selected from the group comprising,
  - (i) a nucleic acid represented by any of SEQ ID NO: 1, 5 or 7 the complement strand thereof;
  - (ii) a nucleic acid encoding an amino acid sequence represented by SEQ ID NO: 2,
     4, 6, 8 or 10 or a homologue, derivative or active fragment of the above mentioned sequences;
  - (iii) a nucleic acid capable of hybridising with a nucleic acid of (i) or (ii) above, which hybridising sequence preferably encodes a protein having seedy1 protein activity;
  - (iv) a nucleic acid which is degenerate from any one of the nucleic acids of (i) to (iii) above as a results of the genetic code;
  - (v) a nucleic acid which is an allelic variant of any one of the nucleic acids of (i) to (iv);
  - (vi) a nucleic acid which is an alternative splice variant of any one of the nucleic acids of (i) to (v);
  - (vii) a nucleic acid encoding a protein which has at least 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 60%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% sequence identity to any one or more from the sequences defined in (i) to (vi), which protein preferably encodes a protein having seedy1 activity;

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(viii) a portion of a nucleic acid according to any of (i) to (vii) above, which portio preferably encodes a protein having seedy1 activity.

According to another embodiment of the present invention, there is provided an isolated nucleic acid as mentioned herein above, wherein said isolated nucleic acid is not a nucleic acid selected from the group comprising: the rice cDNA as deposited under Genbani accession number AK063941 (SEQ ID NO 3), a *Medicago* BAC clone deposited as AC144618 or AC139356 or AC144482 or AC135566, the *Arabidopsis* cDNA deposited under AL61572 (SEQ ID NO 11), or the *Zea mays* EST deposited under AY108162 (SEQ ID NO 9). Further the present invention described for the first time a novel type of protein and therefore, within the scope of the present invention is an isolated seedy1 protein comprising in the following order from N-terminus to C-terminus at least two motifs, which motifs are:

- (i) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 15; and/or
- (ii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 16, and/or
- (iii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 17 and which is a coiled coil motif; and/or
- (iv) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 18.

Further, the present invention provides, an isolated seedy1 protein, comprising

- a. a polypeptide with an amino acid sequence as presented in any one of SEQ ID NO
   2, 4, 6, 8 or 10;
- b. a polypeptide with an amino acid sequence which has at least 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 60%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% sequence identity to any one or more of the amino acid sequence as described in (a)
- 30 c. a polypeptide which is a homologue, a derivative, an immunologically active and/or functional fragment of a protein as defined in any of (a) or (b).

According to another embodiment of the present invention, there is provided an isolated seedy1 protein as mentioned herein above, wherein said protein is not the *Arabisopsis* protein as deposited in Genbank under the accession number AL161572 (SEQ ID NO 12).

According to a preferred aspect of the present invention, enhanced or increased expression of a seedy1 nucleic acid in a plant or plant part is envisaged. Methods for obtaining increased expression of genes or gene products are well documented in the art and include, for example, overexpression driven by a (strong) promoter, the use of transcription enhancers or translation enhancers. The term overexpression as used herein means any form of expression that is additional to the original wild-type expression level. Preferably the nucleic acid to be introduced into the plant and/or the nucleic acid that is to be overexpressed in the plants is in the sense direction with respect to the promoter to which it is operably linked. Alternatively, selection of better performing alleles of the wild-type seedy1 nucleic acid can be achieved via plant breeding techniques.

Accordingly, a preferred embodiment of the present invention provides a method to modify growthe characteristics in a plant, comprising introducing, into a plant, a nucleic acid capable of modifying expression of a seedy1 gene and/or capable of modifying activity and/or level of a seedy1 protein in the sense orientation relative to control element to which it is operably linked.

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Alternatively and/or additionally, increased expression of a seedy1 encoding gene or increased activities and/or levels of a seedy1 protein in a plant cell, is achieved by mutagenesis. For example these mutations can be responsible for the changed control of the seedy1 gene, resulting in more expression of the gene, relative to the wild-type gene. Mutations can also cause conformational changes in a protein, resulting in more activity and/or levels of the seedy1 protein.

Additionally, these is envisaged by the present invention a method for modifying growth characteristics comprising downregulation of expression of a seedy1 gene or downregulation of levels and/or activity of a seedy1 protein. These methods are particularly useful to modify the size or quantity or architecture of plants particular plant organs. According to a particular embodiment of the present invention, plants are decreased in size or plant organs are decreased in size or number, or the plant architecture is changed, for example there is less branching or less branching of the inflorescence. Therefore, according to a further aspect of the invention, decreased expression of a seedy1 nucleic acid or decreased activity and/or level of a seedy1 is envisaged.

Examples of decreasing or downregulation of expression are well documented in the art and include, for example, downregulation of expression by anti-sense techniques, RNAi techniques, small interference RNAs (siRNAs), microRNA (miRNA), etc. Therefore according to a particular aspect of the invention, there is provided a method for modifying growth characteristics of plants, including technologies that are based on for example the synthesis of antisense transcripts, complementary to the mRNA of a seedy1 gene.

Another method for downregulation of gene expression or gene silencing comprises use o ribozymes, for example as described in WO9400012 (Atkins et al.), WO9503404 (Lenee et al.) WO0000619 (Nikolau et al.), WO9713865 (Ulvskov et al.) and WO9738116 (Scott et al.).

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Gene silencing may also be achieved by insertion mutagenesis (for example, T-DNA insertion or transposon insertion) or by gene silencing strategies as described among others in the documents WO9836083 (Baulcombe and Angell), WO9853083 (Grierson et al.), WO9915682 (Baulcombe et al.) or WO9953050 (Waterhouse et al.).

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Expression of an endogenous gene may also be reduced if the endogenous gene contains a mutation. Such a mutant gene may be isolated and introduced into the same or different plant species in order to obtain plants having modified growth characteristics. Also dominant negative mutants of a seedy1 nucleic acid can be introduced in the cell to decrease the level/and or activity of the endogenous Seedy1 protein.

Other methods to decrease the expression of a seedy1 nucleic acid and/or activity and/or level of seedy1 proteins in a cell encompass for example the mechanisms of transcriptional gene silencing, such as the methylation of the seedy1 promoter.

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Another mechanism to downregulate levels and/or activity of a seedy1 protein in a plant encompasses the mechanism of co-suppression. Modifying gene expression (whether by a direct or indirect approach) encompasses changed transcript levels of that gene. Changed transcript levels can be sufficient to induce certain phenotypic effects, for example via the mechanism of cosuppression. Here the overall effect of expression of a transgene is that there is less activity in the cell of the protein encoded by a native gene having homology to the introduced transgene. Cosuppression is accomplished by the addition of coding sequences or parts thereof in a sense orientation into the cell. Therefore, according to one aspect of the present invention, the development of a plant may be changed by introducing into a plant an additional copy (in full or in part) of a seedy1 gene already present in a host plant. The additional gene may silence the endogenous gene, giving rise to a phenomenon known as cosuppression.

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Genetic constructs aimed at silencing gene expression may comprise the seedy1 nucleotide sequence or one at least a portion thereof in a sense and/or antisense orientation relative to the promoter sequence. Preferably the portions comprises at least 21 contiguous nucleic acid of a sequence to be downregulated. Also, sense or antisense copies of at least part of the

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endogenous gene in the form of direct or inverted repeats may be utilised in the methods according to the invention. The development of plants may also be changed by introducing into a plant at least part of an antisense version of the nucleotide sequence represented, for example, by SEQ ID NO: 1. It should be clear that part of the nucleic acid (a portion) could also achieve the desired result. Homologous anti-sense genes are preferred, homologous genes being plant genes, preferably plant genes from the same plant species in which the silencing construct is introduced.

The expression of a seedy1 gene can be investigated by Northern or Southern blot analysis of cell extracts. The levels of seedy1 protein in the cell can be investigated via Western blot analysis of cell extracts.

According to a further embodiment of the present invention, genetic constructs and vectors to facilitate introduction and/or expression of the nucleotide sequences useful in the methods according to the invention are provided. Therefore, according to the further embodiment, the present invention provides a genetic construct comprising:

- (i) a at least part of a nucleic acid of the present invention as described above or at least a part of a nucleic acid encoding a seedy1 protein as defined above;
- (ii) one or more control sequences capable of regulating expression of the nucleic acid of (i); and optionally
- (iii) a transcription termination sequence.

According to methods of the present invention, such a genetic construct is introduced into a plant or plant part.

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Constructs useful in the methods according to the present invention may be constructed using recombinant DNA technology well known to persons skilled in the art. The gene constructs may be inserted into vectors, which may be commercially available, suitable for transforming into plants and suitable for expression of the gene of interest in the transformed cells.

The genetic construct can be an expression vector wherein said nucleic acid is operably linked to one or more control sequences allowing expression in prokaryotic and/or eukaryotic host cells.

The nucleic acid according to (i) is advantageously any of the aforementioned nucleic acids, preferably any of the above mentioned variants of a seedy1 nucleic acid, most preferably a seedy1 nucleic acid, such as for example one of the nucleic acids of SEQ ID NO 1, 3, 5, 7, 9 or

11. The construct sequence of (ii) is preferably a seed-preferred promoter, for example a prolamin promoter

According to a preferred embodiment of the invention, the genetic construct is an expression vector designed to overexpress the nucleic acid. A preferred nucleic acid (i) in this overexpression vector is the sequence represented by SEQ ID NO 1, 3, 5, 7, 9 or 11 or a portion thereof or sequences capable of hybridising therewith or a nucleic acid encoding a sequence represented by SEQ ID NO 2, 4, 6, 8, 10 or 12 or a homologue, derivative or active fragment thereof. Preferably, this nucleic acid is cloned in the sense orientation relative to the control sequence.

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The methods according to the present invention may also be practised by introducing into a plant at least a part of a (natural or artificial) chromosome (such as a <u>Bacterial Artificial Chromosome</u> (BAC)), which chromosome contains at least a seedy1 nucleic acid, optionally together with one or more related gene family members. Therefore, according to a further aspect of the present invention, there is provided a method for modifying growth characteristics of a plant by introducing into a plant at least a part of a chromosome comprising at least a seedy1 gene/nucleic, which seedy1 nucleic is preferably one represented by any one of SEQ ID NO 1, 3, 5, 7, 9 or 11.

Plants are transformed with a vector comprising the sequence of interest (i.e., the nucleic acid 20 capable of modifying expression of seedy1 nucleic acid), which sequence is operably linked to one or more control sequences (at least a promoter). The terms "regulatory element", "control sequence" are all used herein interchangeably and are to be taken in a broad context to refer to regulatory nucleic acids capable of effecting expression of the sequences to which they are ligated (i.e. operably linked). Encompassed by the aforementioned terms are promoters. A 25 "Promoter" encompasses transcriptional regulatory sequences derived from a classical eukaryotic genomic gene (including the TATA box which is required for accurate transcription initiation, with or without a CCAAT box sequence) and additional regulatory elements (i.e. upstream activating sequences, enhancers and silencers) which alter gene expression in response to developmental and/or external stimuli, or in a tissue-specific manner. Also included within the term is a transcriptional regulatory sequence of a classical prokaryotic gene, in which case it may include a -35 box sequence and/or -10 box transcriptional regulatory sequences. The term "regulatory element" also encompasses a synthetic fusion molecule or derivative which confers, activates or enhances expression of a nucleic acid molecule in a cell, tissue or organ. The term "operably linked" as used herein refers to a 35 functional linkage between the promoter sequence and the gene of interest, such that the promoter sequence is able to initiate transcription of the gene of interest.

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Advantageously, any type of promoter may be used to drive expression of the nucleic acid depending on the desired outcome. For example, a meristem-specific promoter, such as the rnr (ribonucleotide reductase), cdc2a promoter and the cyc07 promoter. Also seed -specific promoter, such as p2S2, pPROLAMIN, pOLEOSIN could be selected. An aleurone-specific promoter may be selected. An inflorescence-specific promoter, such as pLEAFY, may also be utilised. To produce male-sterile plants one would need an anther specific promoter. One could also choose a petal-specific promoter. If the desired outcome would be to modify plant growth characteristics in particular organs, then the choice of the promoter would depend on the organ to be changed. For example, use of a root-specific promoter would lead to phenotypic alteration of the root. This would be particularly important where it is the root itself that is the desired end product; such crops include sugar beet, turnip, carrot, and potato. A fruit-specific promoter may be used to modify, for example, the strength of the outer skin of the fruit or to increase the size of the fruit. A green tissue-specific promoter may be used to influence the phenotype pf the leaf. A cell wall-specific promoter may be used to increase the rigidity of the cell wall, thereby increasing resistance of the cell. An anther-specific promoter may be used to produce male-sterile plants. A vascular-specific promoter may be used to increase transport from leaves to seeds. A nodule-specific promoter may be used to increase the nitrogen fixing capabilities of a plant, thereby increasing the nutrient levels in a plant. A stress-inducible promoter may also be used to drive expression of a nucleic acid during conditions of stress. A 20 stress inducible promoter such as the water stress induced promoter WSI18, the drought stress induced Trg-31 promoter, the ABA related promoter rab21 or any other promoter which is induced under a particular stress condition such as temperature stress (cold, freezing, heat) or osmotic stress, or drought stress or oxidative stress or biotic stress can be used to drive expression of a seedy1 gene. 25

Preferably, the nucleic acid capable of modifying expression of a seedy1 gene is operably linked to a plant promoter, preferably a tissue preferred promoter. The term "tissue-preferred" as defined herein refers to a promoter that is expressed predominantly in at least one tissue or organ. Preferably, the tissue-preferred promoter is a seed-preferred promoter or a seedspecific promoter, further preferably an endosperm-specific promoter, more preferably a promoter isolated from a gene encoding a seed-storage protein most preferably a promoter isolated from a prolamin gene, such as for example a rice prolamin promoter as presented in SEQ ID NO 14 or a promoter of similar strength and/or a promoter with a similar expression pattern as the rice prolamin promoter. Similar strength and/or similar expression pattern can be analysed for example by coupling the promoters to a reporter gene and check the function of the reporter gene in tissues of the plant. One suitable reporter gene is beta-glucuronidase and

the colorimetric GUS staining to visualize the beta-glucuronidase activity in a plant tissue i well known to a person skilled in the art.

Examples of preferred seed-specific promoters and other tissue-specific promoters are presented in Table I, which promoters or derivatives thereof are useful in performing the methods of the present invention. Accordingly, genetic constructs comprising at least part of a seedy1 nucleic acid and at least part of a promoter from table I, preferably, wherein said parts are operably linked, is also provided by the present invention.

TABLE I

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THE PRESENT INVE	VIION	
GENE SOURCE	EXPRESSION PATTERN	REFERENCE
seed-specific genes	seed	Simon, et al., Plant Mol. Biol. 5: 19 1985; Scofield, et al., J. Biol. Chel 262: 12202, 1987.; Baszczynski, et a Plant Mol. Biol. 14: 633, 1990.
Brazil Nut albumin	seed	Pearson, et al., Plant Mol. Biol. 18: 23: 245, 1992.
legumin	seed	Ellis, et al., Plant Mol. Biol. 10: 203-214
glutelin (rice)	seed	Takaiwa, et al., Mol. Gen. Genet. 208 15-22, 1986; Takaiwa, et al., FEBS Letts. 221: 43-47, 1987.
zein	seed	Matzke et al Plant Mol Biol, 14(3):323 32 1990
napA	seed	Stalberg, et al, Planta 199: 515 -519 1996.
wheat LMW and HMW glutenin-1	endosperm	Mol Gen Genet 216:81-90, 1989; NAR 17:461-2, 1989
wheat SPA	seed	Albani et al, Plant Cell, 9: 171-184, 1997
wheat α, β, γ-gliadins	endosperm	EMBO 3:1409-15, 1984
barley Itr1 promoter	endosperm	
barley B1, C, D, hordein	endosperm	Theor Appl Gen 98:1253-62, 1999; Plant J 4:343-55, 1993; Mol Gen Genet 250:750-60, 1996
barley DOF	endosperm	Mena et al, The Plant Journal, 116(1): 53-62, 1998
blz2	endosperm	EP99106056.7

, 000		Vicente-Carbajosa et al., Plant J. 13:
synthetic promoter	endosperm	:
		629-640, 1998.  Wu et al, Plant Cell Physiology 39(8)
rice prolamin NRP33	endosperm	885-889, 1998
		Wu et al, Plant Cell Physiology 39(8)
rice α-globulin Glb-1	endosperm	
		885-889, 1998 Sato et al, Proc. Natl. Acad. Sci. USA,
rice OSH1	embryo	
		93: 8117-8122, 1996  Nakase et al. Plant Mol. Biol. 33: 513-
rice α-globulin	endosperm	
REB/OHP-1		522, 1997
rice ADP-glucose PP	endosperm	Trans Res 6:157-68, 1997
maize ESR gene family	endosperm	Plant J 12:235-46, 1997
sorgum γ-kafirin	endosperm	PMB 32:1029-35, 1996
KNOX	embryo	Postma-Haarsma et al, Plant Mol. Biol.
1		39:257-71, 1999
rice oleosin	embryo and aleuron	Wu et at, J. Biochem., 123:386, 1998
sunflower oleosin	seed (embryo and dry seed)	Cummins, et al., Plant Mol. Biol. 19:
		873-876, 1992
Metallothionein Mte, PRO0001		transfer layer of embryo + calli
putative beta-amylase, PRO0005		transfer layer of embryo
putative cellulose synthase, PRO0009		weak in roots
ipase (putative), PRO0012		
transferase (putative), PR		
pentidyl prolyl cis-trans isc	omerase (putative), PRO0016	
Unknown, PRO0019		
prp protein (putative), PRO0020		
noduline (putative), PRO0		
proteinase inhibitor Rgpi9, PRO0058		seed
beta expansine EXPB9, PRO0061		weak in young flowers
structural protein, PRO0063		young tissues+calli+embryo
xylosidase (putative), PR		
prolamine 10 Kda, PRO0075		strong in endosperm
allergen RA2, PRO0076		strong in endosperm
		strong in endosperm
1 -	7	
prolamine RP7, PRO007	7	
prolamine RP7, PRO007 CBP80, PRO0078		
prolamine RP7, PRO007 CBP80, PRO0078 starch branching enzyme	e I, PRO0079	transfer layer of embryo + calli
prolamine RP7, PRO007 CBP80, PRO0078 starch branching enzyme Metallothioneine-like ML:	e I, PRO0079	

prolamine RP6, PRO0090	strong endosperm
prolamine RP5, PRO0091	strong in endosperm
allergen RA5, PRO0092	
putative methionine aminopeptidase, PRO0095	embryo
ras-related GTP binding protein, PRO0098	
beta expansine EXPB1, PRO0104	
Glycine rich protein, PRO0105	· · · · · · · · · · · · · · · · · · ·
metallothionein like protein (putative), PRO0108	
metallothioneine (putative), PRO0109	
RCc3, PR00110	strong root
uclacyanin 3-like protein, PRO0111	weak discrimination center / shoot
	meristem
26S proteasome regulatory particle non-ATPase subunit 11,	very weak meristem specific
PRO0116	
putative 40S ribosomal protein, PRO0117	weak in endosperm
chlorophyll a/b-binding protein presursor (Cab27), PRO0122	very weak in shoot
putative protochlorophyllide reductase, PRO0123	strong leaves
metallothionein RiCMT, PRO0126	strong discrimination center / shoot
	meristem
GOS2, PRO0129	strong constitutive
GOS9, PRO0131	
chitinase Cht-3, PRO0133	very weak meristem specific
alpha-globulin, PRO0135	strong in endosperm
alanine aminotransferase, PRO0136	weak in endosperm
cyclin A2, PRO0138	
Cyclin D2, PRO0139	
Cyclin D3, PRO0140	
cyclophyllin 2, PRO0141	shoot and seed
sucrose synthase SS1 (barley), PRO0146	medium constitutive
trypsin inhibitor ITR1 (barley), PRO0147	weak in endosperm
ubiquitine 2 with intron, PRO0149	strong constitutive
WSI18, PRO0151-	embryo + stress
HVA22 homologue (putative), PRO0156	
EL2, PR00157	
Aquaporine, PRO0169	medium constitutive in young plants
High mobility group protein, PRO0170	strong constitutive
reversibly glycosylated protein RGP1, PRO0171	weak constitutive
cytosolic MDH, PRO0173	shoot
RAB21, PRO0175	embryo + stress
L <del> </del>	

CDPK7, PRO0176	
Cdc2-1, PRO0177	very weak in meristem
sucrose synthase 3, PRO0197	
OsVP1, PRO0198	
OSH1, PRO0200	very weak in young plant meristem
putative chlorophyllase, PRO0208	
OsNRT1, PRO0210	
EXP3, PRO0211	
phosphate transporter OjPT1, PRO0216	
oleosin 18kd, PRO0218	aleurone + embryo
ubiquitine 2 without intron, PRO0219	
RFL, PRO0220	
maize UBI delta intron, PRO0221	
glutelin-1, PRO0223	
fragment of prolamin RP6 promoter, PRO0224	
4xABRE, PRO0225	
glutelin OSGLUA3, PRO0226	
BLZ-2_short (barley), PRO0227	
BLZ-2_long (barley), PRO0228	

Optionally, one or more terminator sequences may also be used in the construct introduced into a plant. The term "terminator" encompasses a control sequence which is a DNA sequence at the end of a transcriptional unit which signals 3' processing and polyadenylation of a primary transcript and termination of transcription. Additional regulatory elements may include transcriptional as well as translational enhancers. Those skilled in the art will be aware of terminator and enhancer sequences, which may be suitable for use in performing the invention. Such sequences would be known or may readily be obtained by a person skilled in the art.

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The genetic constructs of the invention may further include an origin of replication sequence which is required for maintenance and/or replication in a specific cell type. One example is when a genetic construct is required to be maintained in a bacterial cell as an episomal genetic element (e.g. plasmid or cosmid molecule). Preferred origins of replication include, but are not limited to, the f1-ori and colE1.

The genetic construct may optionally comprise a selectable marker gene. As used herein, the term "selectable marker gene" includes any gene, which confers a phenotype on a cell in which it is expressed to facilitate the identification and/or selection of cells which are

transfected or transformed with a nucleic acid construct of the invention. Suitable markers may be selected from markers that confer antibiotic or herbicide resistance, that introduce a new metabolic trait or that allow visual selection. Examples of selectable marker genes include genes conferring resistance to antibiotics (such as nptll encoding neomycin phosphotransferase capable of phosphorylating neomycin and kanamycin, or hpt encoding hygromycin phosphotransferase capable of phosphorylating hygromycin), to herbicides (for example bar which provides resistance to Basta; aroA or gox providing resistance against glyphosate), or genes that provide a metabolic trait (such as manA that allows plants to use mannose as sole carbon source). Visual marker genes result in the formation of colour (for example beta-glucuronidase, GUS), luminescence (such as luciferase) or fluorescence (Green Fluorescent Protein, GFP, and derivatives thereof). Further examples of suitable selectable marker genes include the ampicillin resistance (Ampr), tetracycline resistance gene (Tcr), bacterial kanamycin resistance gene (Kanr), phosphinothricin resistance gene, and the chloramphenical acetyltransferase (CAT) gene, amongst others

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In a preferred embodiment, the genetic construct as mentioned above, comprises at least part of a rice prolamin promoter operably linked to a seedy1 nucleic acid in the sense orientation. An example of such an expression cassette, further comprising a terminator sequence s presented in SEQ ID NO 13.

Therefore the present invention provides an isolated genetic construct, comprising a nucleic acid selected from the group comprising:

- a. A nucleic acid having at least a part of a seed-preferred promoter and at least a part of the nucleic acid as defined herein above 2; and
- b. A nucleic acid as presented in SEQ ID NO 13, or the complementary strand thereof;
- c. a nucleic acid which is degenerated as a result of the genetic code from any of the nucleic acids of (a) or (b);
  - d. a nucleic acids which is an allelic variant of any of the nucleic acids of (a) or (b);
  - e. a nucleic acid which hybridizes to any of the nucleic acids of (a) or (b).
- According to a further embodiment of the present invention, there is provided a method for the production of a plant having modified growth characteristics, comprising modifying expression and or activity and/or levels in a plant of a seedy1 nucleic acid or seedy1 protein.

  According to a particular embodiment, the present invention provides a method for the production of a transgenic plant having modified growth characteristics, which method comprises:

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- (i) introducing into a plant or plant part a nucleic acid capable of modifying expression of a seedy1 gene and/or capable of modifying the activity and/or levels of a seedy1 protein;
- (ii) cultivating the plant cell under conditions promoting regeneration and mature plant growth.

The nucleic acid of (i) may advantageously be any of the aforementioned nucleic acids, preferably a nucleic acid encoding a coiled coil protein, more preferably encoding a seedy1 most preferably a seedy1 nucleic acid according to SEQ ID NO 1, 3, 5, 7, 9, or 11.

Preferably in the above mentioned method, the nucleic acid of (i) is overexpressed in the plant, is in the sense direction and/or is driven by a seed-preferred promoter, such as the prolamin promoter.

The protein itself and/or the nucleic acid itself may be introduced directly into a plant cell or into the plant itself (including introduction into a tissue, organ or any other part of the plant). According to a preferred feature of the present invention, the nucleic acid is preferably introduced into a plant by transformation.

The term "transformation" as referred to herein encompasses the transfer of an exogenous polynucleotide into a host cell, irrespective of the method used for transfer. Plant tissue capable of subsequent clonal propagation, whether by organogenesis or embryogenesis, may be transformed with a genetic construct of the present invention and a whole plant regenerated therefrom. The particular tissue chosen will vary depending on the clonal propagation systems available for, and best suited to, the particular species being transformed. Exemplary tissue targets include leaf disks, pollen, embryos, cotyledons, hypocotyls, megagametophytes, callus tissue, existing meristematic tissue (e.g., apical meristem, axillary buds, and root meristems), and induced meristem tissue (e.g. cotyledon meristem and hypocotyl meristem). The polynucleotide may be transiently or stably introduced into a host cell and may be maintained non-integrated, for example, as a plasmid. Alternatively and preferably, the transgene may be stably integrated into the host genome. The resulting transformed plant cell can then be used to regenerate a transformed plant in a manner known to persons skilled in the art.

Transformation of a plant species is now a fairly routine technique. Advantageously, any of several transformation methods may be used to introduce the gene of interest into a suitable ancestor cell. Transformation methods include the use of liposomes, electroporation, chemicals that increase free DNA uptake, injection of the DNA directly into the plant, particle gun bombardment, transformation using viruses or pollen and microprojection. Methods may be selected from the calcium/polyethylene glycol method for protoplasts (Krens, F.A. et al.,

1882, Nature 296, 72-74; Negrutiu I. et al., June 1987, Plant Mol. Biol. 8, 363-373 electroporation of protoplasts (Shillito R.D. et al., 1985 Bio/Technol 3, 1099-1102 microinjection into plant material (Crossway A. et al., 1986, Mol. Gen Genet 202, 179-185) DNA or RNA-coated particle bombardment (Klein T.M. et al., 1987, Nature 327, 70) infection with (non-integrative) viruses and the like.

Transgenic rice plants expressing a seedy1 gene are preferably produced via *Agrobacterium* mediated transformation using any of the well known methods for rice transformation, such as described in any of the following: published European patent application EP 1198985 A1 Aldemita and Hodges (Planta, 199, 612-617, 1996); Chan *et al.* (Plant Mol. Biol. 22 (3) 491-506, 1993), Hiei *et al.* (Plant J. 6 (2) 271-282, 1994), which disclosures are incorporated by reference herein as if fully set forth. In the case of corn transformation, the preferred method is as described in either Ishida *et al.* (Nat. Biotechnol. 1996 Jun; 14(6): 745-50) or Frame *et al.* (Plant Physiol. 2002 May; 129(1): 13-22), which disclosures are incorporated by reference herein as if fully set forth.

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Generally after transformation, plant cells or cell groupings are selected for the presence of one or more markers which are encoded by plant-expressible genes co-transferred with the gene of interest, following which the transformed material is regenerated into a whole plant.

Following DNA transfer and regeneration, putatively transformed plants may be evaluated, for instance using Southern analysis, for the presence of the gene of interest, copy number and/or genomic organisation. Alternatively or additionally, expression levels of the newly introduced DNA may be monitored using Northern and/or Western analysis, both techniques being well known to persons having ordinary skill in the art.

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The generated transformed plants may be propagated by a variety of means, such as by clonal propagation or classical breeding techniques. For example, a first generation (or T1) transformed plant may be selfed to give homozygous second generation (or T2) transformants, and the T2 plants further propagated through classical breeding techniques.

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The generated transformed organisms may take a variety of forms. For example, they may be chimeras of transformed cells and non-transformed cells; clonal transformants (e.g., all cells transformed to contain the expression cassette); grafts of transformed and untransformed tissues (e.g., in plants, a transformed rootstock grafted to an untransformed scion).

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The present invention also encompasses plants obtainable by the methods according to the present invention. The present invention therefore provides plants obtainable by the method

according to the present invention, which plants have modified growth characteristocs, when compared to the wild-type plants.

The present invention clearly extends to any plant cell or plant produced by any of the methods described herein, and to all plant parts and propagules thereof. The present invention extends further to encompass the progeny of a primary transformed or transfected cell, tissue, organ or whole plant that has been produced by any of the aforementioned methods, the only requirement being that progeny exhibit the same genotypic and/or phenotypic characteristic(s) as those produced in the parent by the methods according to the invention i.e. having modified growth characteristics. The invention accordingly also includes host cells having modified expression of a nucleic acid according to the invention and described herein before and/or having modified activity and/or level of a seedy1 protein or a protein according to the present invention, wherein said expression, level or activity is modified relative to corresponding wild type host cells. Preferred host cells according to the invention are plant cells or cells from insects, animals, yeast, fungi, algae or bacteria. The invention also extends to harvestable parts of a plant such as but not limited to seeds, flowers, stamen, leaves, petals, fruits, stem, stem cultures, rhizomes, roots, tubers and bulbs.

Preferably said host cells is transformed with a seedy1 encoding gene or a genetic construct as described above wherein the seedy1 nucleic acid is under the control of a seed-preferred promoter and more preferably the plants of the present invention carry an expression cassette comprising at least a part of seedy1 and at least a part of a constitutive promoter as mentioned hereinabove. The host cells, plants or the plant parts of the present invention can be identified by the presence of higher expression of a seedy1 gene and/or or a higher level and/or activity of a seedy1 protein. Further, particular plants of the present invention are recognizable by the presence of a seedy1 transgene.

The term "plant" as used herein encompasses whole plants, ancestors and progeny of the plants and plant parts, including seeds, shoots, stems, roots (including tubers), and plant cells, tissues and organs. The term "plant" also therefore encompasses suspension cultures, embryos, meristematic regions, callus tissue, leaves, gametophytes, sporophytes, pollen, and microspores. Plants that are particularly useful in the methods of the invention include all plants which belong to the superfamily *Viridiplantae*, in particular monocotyledonous and dicotyledonous plants including a fodder or forage legume, ornamental plant, food crop, tree, or shrub selected from the list comprising *Acacia spp.*, *Acer spp.*, *Actinidia spp.*, *Aesculus spp.*, *Agathis australis*, *Albizia amara*, *Alsophila tricolor*, *Andropogon spp.*, *Arachis spp*, *Areca catechu*, *Astelia fragrans*, *Astragalus cicer*, *Baikiaea plurijuga*, *Betula spp.*, *Brassica spp.*,

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Bruguiera gymnorrhiza, Burkea africana, Butea frondosa, Cadaba farinosa, Calliandra sp. Camellia sinensis, Canna indica, Capsicum spp., Cassia spp., Centroema pubescent Chaenomeles spp., Cinnamomum cassia, Coffea arabica, Colophospermum mopant Coronillia varia, Cotoneaster serotina, Crataegus spp., Cucumis spp., Cupressus spp Cyathea dealbata, Cydonia oblonga, Cryptomeria japonica, Cymbopogon spp., Cynthe dealbata, Cydonia oblonga, Dalbergia monetaria, Davallia divaricata, Desmodium spp Dicksonia squarosa, Diheteropogon amplectens, Dioclea spp, Dolichos spp., Dorycniui rectum, Echinochloa pyramidalis, Ehrartia spp., Eleusine coracana, Eragrestis spp., Erythrin spp., Eucalyptus spp., Euclea schimperi, Eulalia villosa, Fagopyrum spp., Feijoa sellowiana Fragaria spp., Flemingia spp, Freycinetia banksii, Geranium thunbergii, Ginkgo biloba, Glycin javanica, Gliricidia spp, Gossypium hirsutum, Grevillea spp., Guibourtia coleosperme Hedysarum spp., Hemarthia altissima, Heteropogon contortus, Hordeum vulgare, Hyparrheni rufa, Hypericum erectum, Hyperthelia dissoluta, Indigo incamata, Iris spp., Leptarrhen pyrolifolia, Lespediza spp., Lettuca spp., Leucaena leucocephala, Loudetia simplex, Lotonu bainesii, Lotus spp., Macrotyloma axillare, Malus spp., Manihot esculenta, Medicago sativa Metasequoia glyptostroboides, Musa sapientum, Nicotianum spp., Onobrychis spp. Omithopus spp., Oryza spp., Peltophorum africanum, Pennisetum spp., Persea gratissime Petunia spp., Phaseolus spp., Phoenix canariensis, Phomium cookianum, Photinia spp. Picea glauca, Pinus spp., Pisum sativum, Podocarpus totara, Pogonarthria flecki Pogonarthria squarrosa, Populus spp., Prosopis cineraria, Pseudotsuga menziesi Pterolobium stellatum, Pyrus communis, Quercus spp., Rhaphiolepsis umbellate Rhopalostviis sapida. Rhus natalensis, Ribes grossularia, Ribes spp., Robinia pseudoacacia Rosa spp., Rubus spp., Salix spp., Schyzachyrium sanguineum, Sciadopitys verticillate Sequoia sempervirens, Sequoiadendron giganteum, Sorghum bicolor, Spinacia spp. Sporobolus fimbriatus, Stiburus alopecuroides, Stylosanthos humilis, Tadehagi spp, Taxodiur. distichum, Themeda triandra, Trifolium spp., Triticum spp., Tsuga heterophylla, Vacciniur. spp., Vicia spp., Vitis vinifera, Watsonia pyramidata, Zantedeschia aethiopica, Zea mays amaranth, artichoke, asparagus, broccoli, Brussels sprouts, cabbage, canola, carrol cauliflower, celery, collard greens, flax, kale, lentil, oilseed rape, okra, onion, potato, rice soybean, straw, sugar beet, sugar cane, sunflower, tomato, squash tea, trees. Alternatively algae and other non-Viridiplantae can be used for the methods of the present inventior Preferably the plant according to the present invention is a crop plant selected from rice maize, wheat, barley, millet, oats, rye, soybean, sunflower, canola, sugarcane, alfalfa leguminosae (bean, pea), flax , lupinus, rapeseed, tobacco, tomato, potato, squash, papaya poplar and cotton. Further preferably, the plant according to the present invention is

monocotyledonous plant, most preferably a cereal.

Advantageously, the present invention provides a method for modifying growth characteristics of a plant. According to further embodiments, the invention provides methods for modifying plant growth characteristics, wherein said modified growth characteristics are selected from any one or more of increased yield, increased biomass, modified plant architecture.

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Further preferably, increased yield is increased seed yield en encompasses increased number of filled seeds and/or increased total seed weight per plant.

The term "increased yield" encompasses an increase in biomass in one or more harvestable parts of a plant relative to the total biomass of corresponding wild-type plants. The term also encompasses an increase in seed yield, which includes an increase in the biomass of the seed (seed weight) and/or an increase in the number of (filled) seeds and/or in the size of the seeds and/or an increase in seed volume, each relative to corresponding wild-type plants. An increase in seed size and/or volume may also influence the composition of seeds. An increase in seed yield could be due to an increase in the number and/or size of flowers. An increase in yield might also increase the harvest index, which is expressed as a ratio of the total biomass over the yield of harvestable parts, such as seeds.

The methods of the present invention are used to increase the seed yield of the plant and are therefore particularly favourable to be applied to crop plants, preferably seed crops and cereals. Therefore, the methods of the present invention are particularly useful for plants such as, rapeseed, sunflower, leguminosae (e.g. soybean, pea, bean) flax, lupinus, canola and cereals such as rice, maize, wheat, barley, millet, oats and rye.

Further preferably, increased biomass encompasses increased biomass of above-ground plant tissue, herein determined as above-ground plant area.

Additionally or alternatively, the plants according to the invention have more above ground 25 area. Therefore, the methods of the present invention are additionally and/or alternatively particularly favourable to crops grown for the green tissue and/or grown for the above ground biomass. The methods of the present invention are particularly useful for increasing leaf size and number of grasses and forage crops (such as forage maize, clover, medicago alfalfa etc.). The methods of the present invention are also particularly useful for increasing the stem size of 30

trees (for paper and pulp industry) and sugar cane.

Further preferably, said modified plant architecture encompasses increased number of panicles.

The methods of the present invention clearly change the appearance or morphology of a plant, 35 including any one or more structural features or combination of structural features thereof. Therefore the plants according to the present invention have changed architecture when

compared to the wild-type plants. Other structural features, which may be altered by the methods of the present invention include shape, size, number, position, texture, arrangement and pattern of any cell, tissue or organ or groups of cells, tissues or organs of a plant, including the root, leaf, shoot, stem or tiller, petiole, trichome, flower, inflorescence (financotyledonous and dicotyledonous plants), panicles, petal, stigma, style, stamen, polle ovule, seed, embryo, endosperm, seed coat, aleurone, fibre, cambium, wood, heartwoo parenchyma, aerenchyma, sieve elements, phloem or vascular tissue, amongst others.

The present invention also relates to use of a seedy1 nucleic acid and and/or protein i modifying growth characteristics and to compositions therefore. According to a particular embodiment, the methods of the present invention are used to change growth characteristic of a plant, wherein said growth characteristics is not disease or pathogen resistance, or is not stress tolerance.

The present invention also relates to use of a seedy1 nucleic acid and and/or protein as growth regulator, such as a herbicide or growth stimulator and to compositions therefore.

The present invention also relates to use of a seedy1 nucleic acid and and/or protein as target for an agrochemical compound, such as a herbicide or growth stimulator and 1 compositions therefore.

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According to another aspect of the present invention, advantage may be taken of the nucleotide sequence capable of modifying expression of a seedy1 nucleic acid in breeding programmes. The nucleic acid may be on a chromosome, or a part thereof, comprising at least the seedy1 nucleic acid and preferably also one or more related family members. In a example of such a breeding programme, a DNA marker is identified which may be genetical linked to a gene capable of modifying expression of a seedy1 nucleic acid in a plant, whice gene may be a gene encoding the seedy1 protein itself or any other gene which may direct or indirectly influence expression of the seedy1 gene and/or activity of the seedy1 protein itself. This DNA marker may then be used in breeding programs to select plants having change development.

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Further the use of allelic variants as described herein-above are particular ly useful for conventional breeding programmes, such as in marker-assisted breeding, which is als encompassed by the present invention. Such breeding programmes sometimes require the introduction of allelic variations in the plants by mutagenic treatment of a plant. One suitable mutagenic method is EMS mutagenesis. Identification of allelic variants then may take place by, for example, PCR. Tilling is preferred for identifying allelic variants. This is followed by

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selection step for selection of superior allelic variants of the seedy1 sequence and which give rise to changed development in a plant. Selection, according to the method of the present invention, is typically carried out by monitoring development, differentiation and organ formation of plants containing different allelic variants of the seedy1 sequence, for example, different allelic variants of SEQ ID NO: 1 or of a seedy1 orthologue in that plant. Monitoring growth performance can be done in a greenhouse or in the field. Further optional steps include crossing plants, in which the superior allelic variant was identified, with another plant. This could be used, for example, to make a combination of interesting phenotypic features.

Therefore, mutations in the seedy1 gene may occur naturally, and may form the basis of the selection of plants showing accelerated rate of development, increased organ size and/or number, and/or early flowering.

Accordingly, as another aspect of the invention, there is provided a method for the selection of plants having modified growth characteristics, which method is based on the selection of better-performing allelic variants of the seedy1 sequence relative to the wild-type allele, and which give rise to modified growth characteristics in a plant.

According to a related embodiment of the invention, there is provided a method for breeding plants having modified growth characteristics, comprising the steps of:

- a. Growing a plant with modified expression of a nucleic acid of the present invention as described herein above; and
- b. Crossing the plant of (a) with another plant; and
- c. selecting progeny of the cross of (b) having modified growth characteristic; wherein said growth characteristics are modified relative to the corresponding wild-type plants.
- The cross of step be is for example a cross with a commercial important germplasm.

  Accordingly, the present invention also relates to the use of a seedy1 gene in breeding programmes.

Alternatively, the seedy1 gene itself can be used as a (genetic) marker to detect the presence or absence of a desired trait, or Quantitative Trait Locus (QTLs). In this application of the present invention the gene encoding seedy1 is genetically linked to the desired trait, and typically the phenotypes caused by the gene encoding a seedy1 are monitored in order to breed and select plants with the desired trait. This desired trait or QTL, may comprise a single gene or a cluster of linked genes that affect the desired trait.

In molecular biology it is standard practice to select upon transfection or transformation those individuals (or groups of individuals, such as bacterial or yeast colonies or phage plaques or

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eukaryotic cell clones) that are effectively transfected or transformed with the desired genetic construct. Typically these selection procedures are based on the presence of a selectable of screenable marker in the transfected genetic construct, to distinguish the positive individuals easily from the negative individuals. Therefore, the seedy1 gene can also be used for these purposes, since introduction of this gene into a host cell results in changed development of said host cell.

The methods according to the present invention may also be practised by co-expression of a seedy1 gene in a plant with at least one other gene that cooperates with the seedy1 gene. Co expression may be effected by cloning the genes under the control of a plant expressible promoter in a plant expressible vector and introducing the expression vector(s) into a plant cell using Agrobacterium-mediated plant transformation.

The methods according to the present invention result in plants having modified growth characteristics, as described hereinbefore. These advantageous characteristics may also be combined with other economically advantageous traits, such as further yield-enhancing traits tolerance to various stresses, traits modifying various architectural features and/or biochemica and/or physiological features. Accordingly, the methods of the present invention can also be used in so-called "gene stacking" procedures.

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Also the present invention encompasses a food product derived from any of the plants produced by the methods of the present invention. Further the invention also refers to the use of a product derived from any of the plants according to the present invention in animal feed and in food or in the production procedures thereof.

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In a particular embodiment of the invention the plants with improved growth characteristics resulting in improved yield and/or biomass, are used to produce industrial enzymes and/ or pharmaceuticals. The production of such enzymes or pharmaceuticals in plants is aimed a high accumulation of the desired products in a particular and easy to harvest plant tissues, for example accumulation in the leaves and/or in the seeds. The plants of the present inventior have more seeds and more above ground biomass, and therefore are capable of producing more industrial enzymes and/or pharmaceuticals in these tissues, more particularly in their green biomass and/or in their seeds. Accordingly, the present invention also provides a method for the production of enzymes and/or pharmaceuticals, which method comprises the modifying of expression of a seedy1 gene or the modifying of activity and/or level of a seedy1 protein. Further the invention relates to the use of plants according to the invention for the

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production of industrial enzymes and pharmaceuticals and the invention extends to the industrial enzymes and pharmaceuticals produced according to these methods.

# **Description of the Figures**

5 The present invention will now be described with reference to the following figures in which:

Figure 1 is a schematic presentation of the entry clone, containing CDS0689 within the AttL1 and AttL2 sites for Gateway® cloning in the pDONR201 backbone. CDS0689 is the internal code for the *Nicotiana tabacum* BY2 CDS0689 seedy1 coding sequence. This vector contains also a bacterial kanamycine-resistance cassette and a bacterial origin of replication.

Figure 2 is a map of the binary vector for the expression in *Oryza sativa* of the *Nicotiana tabacum* BY2 seedy1 gene (CDS0689) under the control of the rice prolamin RP6 promoter (PRO0090). This vector contains a T-DNA derived from the Ti Plasmid, limited by a left border (LB repeat, LB Ti C58) and a right border (RB repeat, RB Ti C58)). From the left border to the right border, this T-DNA contains: a selectable marker cassette for antibiotic selection of transformed plants; a screenable marker cassette for visual screening of transformed plants; the PRO0090 - CDS0689 -zein and rbcS-deltaGA double terminator cassette for expression of the *Nicotiana tabacum* BY2 seedy1 gene (CDS0689). This vector also contains an origin of replication from pBR322 for bacterial replication and a selectable marker (Spe/SmeR) for bacterial selection with spectinomycin and streptomycin.

Figure 3 is the alignment of seedy1 nucleic acids and EST's from different plant species. This alignment was made with the program align X of the NVTi software package. The motifs 1, 2, 3 and 4 are indicated with a line.

Figure 4 is the representation of nucleic acids, protein and motif sequences according to the invention.

# 30 Examples

The present invention will now be described with reference to the following examples, which are by way of illustration alone.

Unless otherwise stated, recombinant DNA techniques were performed according to standard protocols described in Sambrook (2001) Molecular Cloning: a laboratory manual, 3rd Edition Cold Spring Harbor Laboratory Press, CSH, New York; or in Volumes 1 and 2 of Ausubel et al.

(1988), Current Protocols in Molecular Biology, Current Protocols. Standard materials at methods for plant molecular work are described in Plant Molecular Biology Labfase (1993) I R.D.D. Croy, published by BIOS Scientific Publications Ltd (UK) and Blackwell Scientific Publications (UK).

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# Example 1: cloning of the seedy1 encoding gene

A cDNA-AFLP experiment was performed on a synchronized tobacco BY2 cell cultu (*Nicotiana tabacum* L. cv. Bright Yellow-2), and BY2 expressed sequence tags that were concycle modulated were identified and elected for further cloning. Subsequently, the expresse sequence tags were used to screen a tobacco cDNA library and to isolate the full-length cDN of interest, namely the cDNA coding for the seedy1 protein of the present invention (CDS0689).

# Synchronization of BY2 cells.

Tobacco BY2 (*Nicotiana tabacum* L. cv. Bright Yellow - 2) cultured cell suspension was synchronized by blocking cells in early S-phase with aphidicolin as follows. Cultured consuspension of *Nicotiana tabacum* L. cv. Bright Yellow 2 were maintained as described (Naga et al. Int. Rev. Cytol. 132, 1-30, 1992). For synchronization, a 7-day-old stationary culture was diluted 10-fold in fresh medium supplemented with aphidicolin (Sigma-Aldrich, St. Louis, MO; mg/l), a DNA-polymerase a inhibiting drug. After 24 h, cells were released from the block to several washings with fresh medium and resumed their cell cycle progression.

#### RNA extraction and cDNA synthesis.

Total RNA was prepared by using LiCl precipitation (Sambrook et al, 2001) and poly(A+) RN was extracted from 500 mg of total RNA using Oligotex columns (Qiagen, Hilden, German according to the manufacturer's instructions. Starting from 1 mg of poly(A+) RNA, first-strar cDNA was synthesized by reverse transcription with a biotinylated oligo-dT25 primer (Gense Paris, France) and Superscript II (Life Technologies, Gaithersburg, MD). Second-strar synthesis was done by strand displacement with *Escherichia coli* ligase (Life Technologies and DNA polymerase I (USB, Cleveland, OH) and RNAse-H (USB).

# cDNA-AFLP analysis.

Five hundred ng of double-stranded cDNA was used for AFLP analysis as described (Vos al., Nucleic Acids Res. 23 (21) 4407-4414, 1995; Bachem et al., Plant J. 9 (5) 745-53, 1996. The restriction enzymes used were BstYl and Msel (Biolabs) and the digestion was done two separate steps. After the first restriction digest with one of the enzymes, the 3' en fragments were collected on Dyna beads (Dynal, Oslo, Norway) by means of their biotinylate

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tail, while the other fragments were washed away. After digestion with the second enzyme, the released restriction fragments were collected and used as templates in the subsequent AFLP steps. For preamplifications, an Msel primer without selective nucleotides was combined with a BstYl primer containing either a T or a C as 3' most nucleotide. PCR conditions were as described (Vos et al., 1995). The obtained amplification mixtures were diluted 600-fold and 5 ml was used for selective amplifications using a P33-labeled BstYl primer and the Amplitaq-Gold polymerase (Roche Diagnostics, Brussels, Belgium). Amplification products were separated on 5% polyacrylamide gels using the Sequigel system (Biorad). Dried gels were exposed to Kodak Biomax films as well as scanned in a phospholmager (Amersham Pharmacia Biotech, Little Chalfont, UK).

#### Characterization of AFLP fragments.

Bands corresponding to differentially expressed transcripts, among which the (partial) transcript corresponding to CDS0689, were isolated from the gel and eluted DNA was reamplified under the same conditions as for selective amplification. Sequence information was obtained either by direct sequencing of the reamplified polymerase chain reaction product with the selective BstYl primer or after cloning the fragments in pGEM-T easy (Promega, Madison, Wl) or sequencing of individual clones. The obtained sequences were compared against nucleotide and protein sequences present in the publicly available databases by BLAST sequence alignments (Altschul et al., Nucleic Acids Res. 25 (17) 3389-3402 1997). When available, tag sequences were replaced with longer EST or isolated cDNA sequences to increase the chance of finding significant homology. The physical cDNA clone corresponding to CDS0689 was subsequently amplified from a commercial Tobacco cDNA library as follows.

# 25 Cloning of a tobacco CDS0689 seedy1 gene (CDS0689)

A c-DNA library with average inserts of 1,400 bp was made with poly(A+) isolated from actively dividing, non-synchronized BY2 tobacco cells. These library-inserts were cloned in the vector pCMVSPORT6.0, comprising an attB gateway cassette (Life Technologies). From this library 46,000 clones were selected, arrayed in 384-well microtiter plates, and subsequently spotted in duplicate on nylon filters. The arrayed clones were screened by using pools of several hundreds of radioactively labelled tags as probe (among which the BY2-tag corresponding to the sequence CDS0689). Positive clones were isolated (among which the clone reacting with the BY2-tag corresponding to the sequence CDS0689), sequenced, and aligned with the tag sequence. Alternatively, when the hybridization with the tag would fail, the full-length cDNA corresponding to the tag was selected by PCR amplification as follows. Tag-specific primers designed was using primer3 program (http://wwwgenome.wi.mit.edu/genome software/other/primer3.html) and used in combination with the

common vector primer to amplify partial cDNA inserts. Pools of DNA from 50.000, 100.00 150.000, and 300.000 cDNA clones were used as templates in the PCR amplification Amplification product were isolated from agarose gels, cloned, sequenced and aligned wittags. The vector comprising the sequence CDS0689 and obtained as described above, we referred to as entry clone.

# Example 2: Vector construction for transformation with PRO0090-CDS068 cassette

The entry clone was subsequently used in an LR reaction with p0830, a destination vector used for *Oryza sativa* transformation. This vector contains as functional elements within the DNA borders: a plant selectable marker; a plant screenable marker; and a Gateway cassett intended for LR in vivo recombination with the sequence of interest already cloned in the enticlone. The rice prolamin RP6 promoter for endosperm-specific expression (PRO0090) located upstream of this Gateway cassette.

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After the LR recombination step, the resulting expression vector as shown in Fig. 2 can b transformed into *Agrobacterium* and subsequently into *Oryza sativa* plants. Transformed ric plants were allowed to grow and then examined for various parameters as described i example 3.

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# Example 3: Evaluation of T0, T1 and T2 transgenic rice plants transformed with prolamin::seedv1 (PRO0090-CDS0689) and results

Approximately 15 to 20 independent T0 rice transformants were generated. The primar transformants were transferred from tissue culture chambers to a greenhouse for growing an harvest of T1 seed. 4 events, of which the T1 progeny segregated 3:1 for presence/absence of the transgene, were retained. For each of these events, approximately 10 T1 seedling containing the transgene (hetero- and homo-zygotes), and approximately 10 T1 seedling lacking the transgene (nullizygotes), were selected by monitoring screenable marks expression.

2 events with improved agronomical parameters in T1 were chosen for re-evaluation in T2 Seed batches from the positive plants (both hetero- and homozygotes) in T1, were screened by monitoring marker expression. For each chosen event, the heterozygote seed batches were then selected for T2 evaluation. An equal number of positives and negatives within each seed batch were transplanted for evaluation in the greenhouse. The total number of 120 seedy transformed plants were evaluated in the T2 generation. More particularly, two seedy transformed events have been selected, 60 plants per event of which 30 positives for the transgene, and 30 negative.

T1 and T2 plants were transferred to the greenhouse and evaluated for vegetative growth parameters and seed parameters, as described hereunder.

Table 1: overview of plants involved in T1 and T2 plant evaluation

Number and types	Number of plants per	Number of positive s	Number of negatives
of lines per evaluation	evaluation	per line	per line
4 lines of T1 plants	80	10	10
2 lines of T2 plants	120	30	30
1 line of T1 plants	20	10	10
1 line of T2 plants	60	30	30

# Statistical analysis: t-test and F-test

A two factor ANOVA (analysis of variants) was used as statistical model for the overall evaluation of plant phenotypic characteristics. An F-test was carried out on all the parameters measured, for all of the plants of all of the events transformed with the gene of interest. The F-test was carried out to check for an effect of the gene over all the transformation events and to determine the overall effect of the gene or "global gene effect". Significant data, as determined by the value of the F-test, indicates a "gene" effect, meaning that the phenotype observed is caused by more than the presence or position of the gene. In the case of the F-test, the threshold for significance for a global gene effect is set at a 5% probability level.

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To check for an effect of the gene within an event, i.e., for a line-specific effect, a t-test was performed within each event using data sets from the transgenic plants and the corresponding null plants. "Null plants" or "Null segregants" are the plants treated in the same way as the transgenic plant, but from which the transgene has segregated. Null plants can also be described as the homozygous negative transformants. The threshold for significance for the t-test is set at 10% probability level. Within one population of transformation events, some events can be under or above this t-test threshold. This is based on the hypothesis that a gene might only have an effect in certain positions in the genome, and that the occurrence of this position-dependent effect is not uncommon. This kind of gene effect may also be referred to as a "line effect of a gene". The p-value is obtained by comparing the t-value to the t-distribution or alternatively, by comparing the F-value to the F-distribution. The p-value stands for the probability that the null hypothesis (null hypothesis being "there is no effect of the transgene") is correct.

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# **Vegetative growth measurements**

The selected transgenic plants were grown in a greenhouse. Each plant received a unique barcode label to link unambiguously the phenotyping data to the corresponding plant. The selected transgenic plants were grown on soil in 10 cm diameter pots under the following environmental settings: photoperiod= 11.5 h, daylight intensity= 30,000 lux or more, daytime temperature= 28°C or higher, night time temperature= 22°C, relative humidity= 60-70%. Transgenic plants and the corresponding nullizygotes were grown side-by-side at randor positions. From the stage of sowing until the stage of maturity each plant was passed severatimes through a digital imaging cabinet and imaged. At each time point digital image (2048x1536 pixels, 16 million colours) were taken of each plant from at least 6 different angles. The parameters described below were derived in an automated way from all the digital image of all the plants, using image analysis software.

## (a) Aboveground plant area

15 Plant aboveground area was determined by counting the total number of pixels from aboveground plant parts discriminated from the background. This value was averaged for the pictures taken on the same time point from the different angles and was converted to a physical surface value expressed in square mm by calibration. Experiments show that the aboveground plant area measured this way correlates with the biomass of plant parts above ground.

# b) Number of primary panicles

The tallest panicle and all the panicles that overlap with the tallest panicles when aligner vertically were counted manually, and considered as primary panicles.

# Seed-related parameter measurements

The mature primary panicles of T1 and T2 plants were harvested, bagged, barcode-labelled and then dried for three days in the oven at 37°C. The panicles were then threshed and all the seeds were collected and counted. The filled husks were separated from the empty one using an air-blowing device. The empty husks were discarded and the remaining fraction was counted again. The filled husks were weighed on an analytical balance. This procedure resulted in the set of seed-related parameters described below.

# (c) Number of filled seeds

35 The number of filled seeds was determined by counting the number of filled husks tha remained after the separation step.

# (d) Total seed yield per plant

The total seed yield was measured by weighing all filled husks harvested from a plant.

The results are given in % of difference between the positive plants and the corresponding nullizygotes (negative) plants of a transgenic line. The values given in table 2 to 4 are the average of 4 T1 lines and the average for T2 lines.

Table 2: overview of phenotypic data of seedy1 transgenic T1 and T2 plants for above ground area

••••••••••••••••••••••••••••••••••••••	% difference betw	reen pos. and neg. plants for above ground	area
	T1 plants	T2 plants	
4 lines	+ 12.5 %		
2 lines		+ 25.5 %	

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Table 3: overview of phenotypic data of seedy1 transgenic T1 and T2 plants for number of first panicles

	% difference betwee	n pos. and neg. plants for nr. of first panicles
	T1 plants	T2 plants
4 lines	+ 32.25 %	
2 lines		+ 26.5 %

# Table 4: overview of phenotypic data of seedy1 transgenic T1 and T2 plants for number of filled seeds

\	% difference between	en pos. and neg. plants for nr. of filled seeds
	T1 plants	T2 plants
4 lines	+ 59 %	
2 lines		+ 36.5 %

Table 5: overview of phenotypic data of seedy1 transgenic T1 and T2 plants for total seed weight per plant

	<u>% difference between po</u>	s. and neg. plants for total seed weight per plant
	T1 plants	T2 plants
4 lines	+70 %	
2 lines		+ 47 %
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# Example 4: Use of the invention in corn

The invention described herein is also be used in maize. To this aim, a seedy1 encoding gent for example a maize or other ortholog, is cloned under control of a seed-specific promote operable in maize, in a plant transformation vector suitable for Agrobacterium-mediated cor transformation. Methods to use for corn transformation have been described in literatur (Ishida et al., Nat Biotechnol. 1996 Jun;14(6):745-50; Frame et al., Plant Physiol. 200 May;129(1):13-22).

10 Transgenic plants made by these methods are grown in the greenhouse for T1 see production. Inheritability and copy number of the transgene are checked by quantitative rea time PCR and Southern blot analysis and expression levels of the transgene are determine by reverse PCR and Northern analysis. Transgenic lines with single copy insertions of th transgene and with varying levels of transgene expression are selected for T2 see production.

Progeny seeds are germinated and grown in the greenhouse in conditions well adapted for maize (16:8 photoperiod, 26-28°C daytime temperature and 22-24°C nighttime temperature as well under water-deficient, nitrogen-deficient, and excess NaCl conditions. Null segregant from the same parental line, as well as wild type plants of the same cultivar are used a controls. The progeny plants resulting from the selfing or the crosses are evaluated on differer biomass and developmental parameters, including, stem size, number of leaves, total above ground area, leaf greenness, time to maturity, flowering time, time to flower, ear number harvesting time. The seeds of these lines are also checked on various parameters, such as grain size, total grain yield per plant, and grain quality (starch content, protein content and o content).

Lines that are most significantly improved compared to corresponding control lines an selected for further field-testing and marker-assisted breeding, with the objective of transferring the field-validated transgenic traits into commercial germplasm. The testing of maize for growth and yield-related parameters in the field is conducted using well-established protocols. Similarly, introgressing specific loci (such as transgene containing loci) from one germplasm into another is also conducted using well-established protocols.

# Claims

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- 1. An isolated nucleic acid encoding at least part of a seedy1 protein, wherein sa seedy1 protein comprises in the following order from N-terminus to C-terminus at least tw motifs selected from:
  - (i) a motif having at least 80% sequence identity to the sequence represented by SE ID NO 15; and/or
  - (ii) a motif having at least 80% sequence identity to the sequence represented by SE ID NO 16, and/or
- 10 (iii) a motif having at least 80% sequence identity to the sequence represented by SE ID NO 17, a coiled coil motif; and/or
  - (iv) a motif having at least 80% sequence identity to the sequence represented by SE ID NO 18.
- 15 2. An isolated nucleic acid selected from:
  - (i) a nucleic acid represented by any of SEQ ID NO: 1, 5, 9, or the complement stranthereof;
  - (ii) a nucleic acid encoding an amino acid sequence represented by SEQ ID NO: 2, 6, 8 or 10 or a homologue, derivative or active fragment of any of the aforementioned sequences;
  - (iii) a nucleic acid capable of hybridising with a nucleic acid of (i) or (ii) above, which hybridising sequence preferably encodes a protein having seedy1 protein activity;
  - (iv) a nucleic acid which is degenerate from any one of the nucleic acids of (i) to (ii) above as a results of the genetic code;
- 25 (v) a nucleic acid which is an allelic variant of any one of the nucleic acids of (i) to (iv);
  - (vi) a nucleic acid which is an alternative splice variant of any one of the nucleic acid of (i) to (v);
- (vii) a nucleic acid encoding a protein which has at least 21%, 22%, 23%, 24%, 25% 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39% 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 60%, 70%, 75% 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% sequence identity to any one comore of the sequences defined in (i) to (vi), which protein preferably encodes protein having seedy1 activity;
- (viii) a portion of a nucleic acid according to any of (i) to (vii) above, which portio preferably encodes a protein having seedy1 activity.

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- 3. An isolated seedy1 protein comprising in the following order from N-terminus to C-terminus at least two motifs selected from:
  - (i) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 15; and/or
- 5 (ii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 16, and/or
  - (iii) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 17, a coiled coil motif; and/or
- (iv) a motif having at least 80% sequence identity to the sequence represented by SEQ ID NO 18.
  - 4. An isolated seedy1 protein, comprising
    - a. a polypeptide with an amino acid sequence as presented in any one of SEQ ID NO
       2, 4, 6, 8 or 10;
- b. a polypeptide with an amino acid sequence which has at least 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 60%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% sequence identity to any one or more of the amino acid sequence as described in (a)
- c. a polypeptide which is a homologue, derivative, immunologically active and/or functional fragment of a protein as defined in any of (a) or (b).
  - An isolated genetic construct comprising:
    - (i) a nucleic acid of claim 1 or 2 or a nucleic acid encoding a protein according to claim 3 or 4;
    - (ii) one or more control sequence capable of regulating expression of the nucleic acid of (i); and optionally
    - (iii) a transcription termination sequence.
- 30 6. An isolated genetic construct according to claim 5, wherein said nucleic acid is represented by SEQ ID NO 1 or a portion thereof or by sequences capable of hybridising therewith, which nucleic acid is preferably from a dicotyledonous plant, further preferably from the family Solanaceae, more preferably the nucleic acid is from Nicotiana, or is a nucleic acid encoding an amino acid sequence represented by SEQ ID NO: 2 or a homologue thereof having at least, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%

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45%, 46%, 47%, 48%, 49%, 50%, 60%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%; 98° or 99% sequence identity with SEQ ID NO 2, or derivative or active fragment thereof.

- 7. An isolated genetic construct according to claim 5 or 6, wherein said control sequenc is a plant promoter, preferably a seed-specific promoter, more preferably an endospern specific promoter, more preferably a promoter isolated from a gene encoding a seet storage protein, most preferably a promoter isolated from a prolamin gene, such as for example the rice prolamin promoter of SEQ ID NO 14.
- 10 8. An isolated genetic construct according to claim 7, comprising an nucleic acid selecte from:
  - a. A nucleic acid having at least a part of a seed-preferred promoter and the nuclei acid as defined in claims 1 or 2; and
  - b. A nucleic acid as presented in SEQ ID NO 13, or the complementary strand thereo
  - a nucleic acid which is degenerated as a result of the genetic code from any of the nucleic acids of (a) or (b);
    - d. a nucleic acids which is an allelic variant of any of the nucleic acids of (a) or (b);
    - e. a nucleic acid which hybridises to any of the nucleic acids of (a) or (b).
- 9. Method for modifying growth characteristics of a plant, comprising modifying in a plant expression of a nucleic acid according to claim 1 or 2 and/or modifying in a plant level and/or activity of a seedy1 protein according to claim 3 or 4, wherein said growth characteristics are modified relative to corresponding wild-type plants.
- 25 10. Method according to claim 9, wherein said modification is effected by recombinar means and/or chemical means.
- Method according to claim 9 or 10, comprising introducing, into a plant, a nucleic acicapable of modifying expression of a gene encoding a seedy1 protein and/or capable c
   modifying levels and/or activity of a seedy1 protein.
  - Method according to claim 11, wherein said nucleic acid is a nucleic acid of claim 1 c2.
- 35 13. Method according to claim 12, wherein said nucleic acid is as represented by SEQ II NO 1 or a portion thereof or sequences capable of hybridising therewith, which nucleic aci is preferably from a dicotyledonous plant, further preferably from the family Solanaceae

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more preferably the nucleic acid is from *Nicotiana*, or is a nucleic acid encoding an amino acid sequence represented by SEQ ID NO 2 or a homologue thereof having at least, 20%, 21%, 22%, 23%, 24%, 25%, 26%, 27%, 28%, 29%, 30%, 31%, 32%, 33%, 34%, 35%, 36%, 37%, 38%, 39%, 40%, 41%, 42%, 43%, 44%, 45%, 46%, 47%, 48%, 49%, 50%, 60%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% sequence identity with SEQ ID NO 2, or derivative or active fragment thereof.

- 14. Method according to any of claims 9 to 13, wherein said nucleic acid is an allelic variant of a nucleic acid encoding the seedy1 protein of claim 2 or 3, or wherein said seedy1 protein is encoded by an allelic variant.
- 15. Method according to any of claims 9 to 13, wherein said nucleic acid is an alternative splice variant of a nucleic acid encoding the seedy1 protein of claim 2 or 3, or wherein said seedy1 protein is encoded by a splice variant.
- 16. Method according to any claims 9 to 13, wherein said nucleic acid is comprised on at least a part of an artificial chromosome, which artificial chromosome preferably also comprises one or more related gene family members.
- 20 17. A method for the production of a transgenic plant having modified growth characteristics, which method comprises:
  - (i) Introducing into a plant cell a nucleic acid of claim 1 or 2;
  - (ii) cultivating the plant cell under conditions promoting regeneration and mature plant growth.
  - 18. Method according to claim 17, wherein said nucleic acid is overexpressed in a plant.
  - 19. Method according to any of claims 17 or 18, wherein said nucleic acid is introduced in the sense direction into a plant.
  - 20. Method according to any of claims 17 to 19, wherein expression of said nucleic acid is driven by a seed-preferred promoter.
- 21. Method according to any of claims 17 to 20, wherein expression of said nucleic acid is driven by a promoter of a seed-storage protein.
  - 22. Method according to claim 21, wherein the promoter is a prolamin promoter.

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- 23. Method for breeding plants having modified growth characteristics, comprising th steps of:
  - (i) Growing a plant with modified expression of a nucleic acid of claim 1 or 2; and
  - (ii) Crossing the plant of (i) with another plant; and
    - (iii) selecting progeny of the cross of (ii) having modified growth characteristics; wherein said growth characteristics are modified relative to the corresponding wild-typ plants.
- 10 24. Method according to any of claims 9 to 23, wherein said modified growth characteristic is selected from any one or more of increased yield, increased biomass, modified plar architecture, each relative to corresponding wild type plants.
- 25. Plants obtainable by a method according to any of claims 9 to 24, which plants hav modified growth characteristics.
  - 26. A host cell having modified expression of a nucleic acid of claim 1 or 2 and/or havin modified activity and/or level of a protein of claim 2 or 4, wherein said expression, level or activity is modified relative to corresponding wild type host cells.
  - 27. A host cell according to claim 26, comprising a nucleic acid of claim 1 or 2 or and/c containing a genetic construct of any one of claims 5 to 8.
- 28. A host cell according to claims 26 or 27, which is a cell from a plant, insect, animal yeast, fungus, algae or bacterium.
  - 29. A plant having modified growth characteristics, comprising the host cell of claim 28.
- 30. A plant according to claim 25 or 29, wherein said plant is a crop plant selected fron rice, maize, wheat, barley, millet, oats, rye, soybean, sunflower, canola, sugarcane, alfalfa leguminosae (bean, pea), flax, lupinus, rapeseed, tobacco, tomato, potato, squash papaya, poplar and cotton, preferably said plant is a monocotyledonous plant, more preferably a cereal.
- 35 31. Plant part, preferably a harvestable plant part, a propagule or progeny from a plan according to claim 25, 29 or 30.

- 32. Use of a nucleic acid of claim 1 or 2 and/or of a seedy1 protein of claim 3 or 4 in modifying the growth characteristics of a plant.
- 33. Use of a nucleic acid of claim 1 or 2 or of a seedy1 protein of claim 3 or 4 as

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- (i) a growth regulator, such as a herbicide or a growth stimulator; or
- (ii) as a target for an agrochemical compound, such as a herbicide or a growth stimulator; or
- (iii) use of a nucleic acid of claim 1 or 2 in breeding programs; or
- 10 (iv) use of a plant according to any one of claims 25, 29 or 30, or use of a product derived therefrom in animal feed or in a food product.
  - (v) Use of plants according to any one of claims 21, 29 or 30 for the production of (industrial) enzymes, pharmaceuticals or plant effective agents.
- 15 34. (Industrial) enzymes, pharmaceuticals and plant effective agents, produced in a plant according to any one of claims 25, 29 or 30.
  - 35. A food product derived from a plant according to any one of claims 21, 29 or 30.
- 20 36. A growth regulating composition, such as a herbicide or a growth stimulator, comprising a nucleic acid of claim 1 or 2 and/or comprising a seedy1 protein of claim 3 or 4, said composition further comprising substances normally added to growth regulating compositions.

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# **Abstract**

# Plants having modified growth characteristics and a method for making the same

The present invention concerns a method for modifying growth characteristics of a plant, in particular increasing biomass and seed yield and modifying plant architecture, by modifying expression of a seedy1 nucleic acid and/or modifying levels and/or activity in a plant of a seedy1 protein. The invention also relates to isolated nucleic acids encoding a seedy1 protein, which seedy1 protein has a coiled coil domain as represented in SEQ ID NO 17 and three conserved motifs as presented in SEQ ID NO 15, 16 and 18. Further the invention relates to transgenic plants having modified growth characteristics, which plants have modified expression of a nucleic acid encoding a seedy1 protein or modified levels and/or activities of a seedy1 protein.

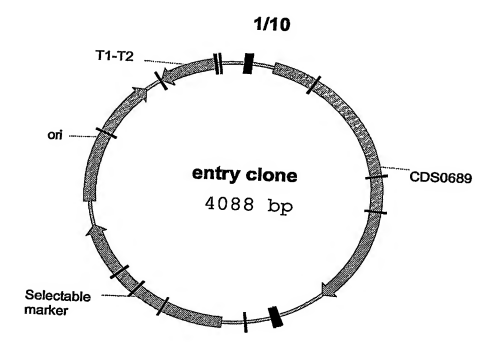


FIGURE 1

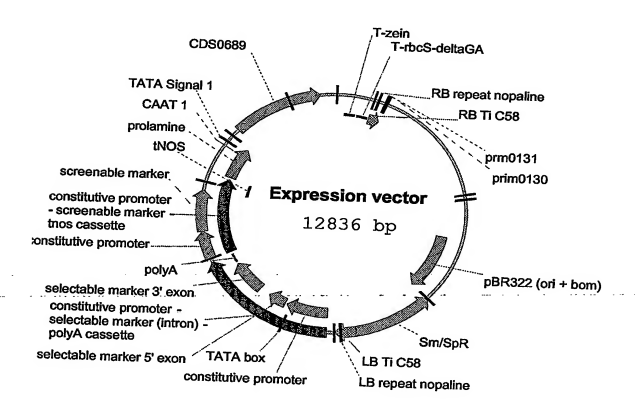


FIGURE 2

		1 Motif 1 40
CDS0689	(1)	msvloypegidpadvoenneafongösedlssäkr
CDS0689 At	(1)	MTSTEATETLNAPPKEQTINNEAFIDGDSQITSATEA
CDS0689 Medicago trunculata	(1)	mnntnnnnillhstqvqvin fafigedfamnsssds
CDS0689 Os	(1)	meedplijplvhynn aafidsscsrsawijpo
CDS0689 Ta variant	(1)	mmeedpiäpivhänäafässsssäwhäha meedpiäpivhänäafähsssawhähsp
CDS0689 So	(1)	meedpi Qivhyisi ac maaissawakase
CDS0689 Zm variantrev CDS0689 So variant	(1) (1)	MEEDPLEPLVHYIN PPT HASYSAWHÄHSP
CDS0689 Bn est1	(1)	mtstehtetlnaperoundaafidgisnijsäjea
CDS0689 Eschscholzia californica	(1)	MLEISETLNEPDEQTVIN AFOSGSEDNHTTAIK
CDS0689 Ga est1	(1)	MSELOYPESFNUPERQUUN AFFINGESEDENARKD
CDS0689 Pt	(1)	mssilqypivvdapivqinitaringisegslniks
CDS0689 Plumbao zeylanica	(1)	MNEVLH QEAARTDSSTUHQUIN AF SGESEDSPVVID
CDS0689 Citrus sinensis	(1)	msyloyphtlngoholwilafingsedettikg I E IPEVQIWNNAAFD GDS S AI
Consensus	(1)	
		41 Motif 2 80
CDS0689	(37)	swsplkplsvrpsdsfesdlsskrightlfenssvnls
CDS0689 At		SSWSHLNESFDSDCSKOPFISVSSSLQSS
CDS0689 Medicago trunculata	(38)	i Lin Safn
CDS0689 Os CDS0689 Ta variant	(32) (33)	TPVRREE REAETN
CDS0689 Ta Variant	(32)	VPASÄRREAEGD ENHREDPDP
CDS0689 Zm variantrev	(32)	spapasarigicol ilreepdv
CDS0689 So variant	(32)	ARASAGHEAEGD HREDPDP
CDS0689 Bn est1	(37)	swsningsfiegcs of QIPVSVSSSLKSS
CDS0689 Eschscholzia californica	(35)	ASSSPLKPIVLNQSEPSILDSIYT QT SCCISPVRTK
CDS0689 Ga est1	(37)	SWCNFNSGSVNQSLBSRGSKDQSELWIKSPVSFK
CDS0689 Pt	(38)	SWWNQSLESDES WLSEVCEQSSPVFV FSAPNLSQELLSDSSI WLSESLAEMPHPAK
CDS0689 Plumbao zeylanica CDS0689 Citrus sinensis	(41) (37)	SWANLKSVYMNQSLESECSE LSERLNKSPTSSL
Consensus	(41)	S ESDG KEN P S .
	•	81 120
CDS0689		SPLPIKPLNPNGALENSRLKPNKPNSKQSLDEMAARKSGK
CDS0689 At	(49)	VSITEAPSSNK-
CDS0689 Medicago trunculata CDS0689 Os	(53)	VAAG
CDS0689 Ta variant	(50)	
CDS0689 So	(54)	
CDS0689 Zm variantrev	(55)	
CDS0689 So variant	(54)	TRACTURE - EVAL
CDS0689 Bn est1	(67)	VSFSTDDPEKPH SPLPIKPLHPNGEKPH
CDS0689 Eschscholzia californica CDS0689 Ga estl	(75) (72)	STASVVKPLSSKNVTGNTREPFSAKMKSGVCKE
CDS0689 Pt	(66)	NSSKPAKPLQ
CDS0689 Plumbao zeylanica	(73)	SPMOK
CDS0689 Citrus sinensis	(72)	KSCVPNKPLQVNSSVKNSQMKQLKSVSK
Consensus	(81)	Motif 3: coiled coil 160
CDS0689	(115)	GNDFRDEKK DEEIEEIOMEISRLSSRLEALF EKAEKTV
CDS0689 At	(94)	RDEDAEIEFÜEKEIGRLSÜKLESLASEKAEQTA
CDS0689 Medicago trunculata	(56)	rtidde iaeïese ikrijišilellriekaerki ydviae ighifacilrisski hhlriskopepn
CDS0689 Os CDS0689 Ta variant	(57)	DADAEIARIEAEILRLSSRLHHLRYSKGHDAK
CDS0689 Id Variant	(54)	DYEAEIGHIEAEILRLXSRLHHLRTSKQSEPS
CDS0689 Zm variantrev	(55)	EEFMRHIEAEILRLSLRLHHLRTSQQLQPP
CDS0689 So variant	(54)	DWEAFIGHIEAFILRLSSRLHHLRTSKQSEPP
CDS0689 Bn est1		ktgkvrhgd daeiee ekemnrlsirieslajekaeqia
CDS0689 Eschscholzia californica	(87)	DOWN DOWN THE THE PERSON DE CARE DECIDED ON THE
CDS0689 Ga est1		eekkrdekkiidmeieeiekeiäarlsikleslriiekpnimq
CDS0689 Ptumbao zeylanica	(76) (78)	
CDS0689 Piumbao zeyianica CDS0689 Citrus sinensis	(3.007	ELETRDERKÄDIE I ELE I EKE I SRI SRLEALRÄEKIDIKT
Consensus	(121)	

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401
                   CDS0689
                           (339) IQSSVVRKRSLPENDKDESKRNDKKRSLSVGKTRVSQTES
                           (294) TGEKDVRKRSLPEDEEKENHKRSEKR-----RASDESN
                CDS0689 At
CDS0689 Medicago trunculata
                            (284) GGD--ARKRSFSEN-----
                           (297) AAAAATAKRMAGSSKMRVIPSRYSÜTPGÄSLÖSSGAQERR
               CDS0689 Os
                           (111) ISTASTCRRPAGSSKVRVVPSRYSIMPG SIG-AATQDGR
(1) -----GRYSIMPG SIGAASQERRR
     CDS0689 Hv contig 123
         CDS0689 Ta contig
                             (87) TSNVATTKRPAGSSKVÄVVPSRYSTPPGSSLÄAVTQGNRC
        CDS0689 Zm partial
        CDS0689 Sacc sp 3'
                            (84)
                                TSNAATAKRPAGSSKVRVVPSRYSITPGSYLAAVSQDKRS
    CDS0689 Pinus taeda 3'
                             (1)
                            (401)
                                                     RYSL PGA LG
                 Consensus
                                 441
                           (379) KNL---GTESRVKKRWEIPSEIVVHGNTESEKSPLJIVK
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CDS0689 Medicago trunculata
                           (337) RKQSLPGSSGDANQNEEIRAKVI PSN--DPLSPOTISKV
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                             (1) -----xearinfgtgnsaimaggtkappitterh
    CDS0689 Pinus taeda 3'
                 Consensus
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                                                      E
                                                              LSPESL KV
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    CDS0689 Pinus taeda 3'
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    CDS0689 Pinus taeda 3'
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                                              ARVLE EA E
                           (521)
                 Consensus
                   CDS0689
                           (476)
                CDS0689 At
                           (403) -----
CDS0689 Medicago trunculata
                           (395) -----
                            (432) -----
                CDS0689 Os
     CDS0689 Hv contig 123
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         CDS0689 Ta contig
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        CDS0689 Zm partial
                            (226) -----
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                           (217)
                             (99) ------
    CDS0689 Pinus taeda 3'
                               Consensus (561)
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# SEQ ID NO 1: Nicotiana tabacum seedy1 coding sequence (CDS0689)

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# SEQ ID NO 2: Nicotiana tabacum seedy1 protein (CDS0689)

MSVLQYPEGIDPADVQIWNNAAFDNGDSEDLSSLKRSWSPLKPLSVRPSDSFESDLSSKENQ TPLFENSSVNLSSPLPIKPLNPNGALENSRLKPNKPNSKQSLDEMAARKSGKGNDFRDEKKI DEEIEEIQMEISRLSSRLEALRIEKAEKTVAKTVEKRGRVVAAKFMEPKQSVIKIEERISMS ARTKVEQRRGLSLGPSEIFTGTRRRGLSMGPSDILAGTTKARQLGKQEMIITPIQPIQNRRK SCFWKLQEIEEEGKSSSLSPKSRKTAARTMVTTRQAVTTIASKKNLKKDDGLLSSVQPKKLF KDLEKSAAANKKPQRPGRVVASRYNQSTIQSSVVRKRSLPENDKDESKRNDKKRSLSVGKTR VSQTESKNLGTESRVKKRWEIPSEIVVHGNTESEKSPLSIIVKPDLLPRIRIARCVNETLRD SGPAKRMIELIGKKSFFSSDEDKEPPVCQVLSFAEEDAEEE

# SEQ ID NO 3: Oryza sativa seedyl coding sequence

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# SEQ ID NO 4: Oryza sativa seedyl protein

MEEDPLIPLVHVWNNAAFDDSSCSRSAWLPQSPAVAAVRKGDKENHRPEVVDVAAGYDVEAE IGHIEAEILRLSSRLHHLRVSKQPEPNRDDAPMGEMVAKVRPRPRGLSLGPLDVISIVNREK HPLRTKQPPATRGRGLSLGPMEIAAANPRVPAAAQHQQQQRAGTARILKPIKEPPVQRRRGV SLGPLEIHHGVGSKAPAAARAKPFTTKLNAIREETRPSKQFAVPAKPWPSSNTRQTLDSRQG TAASRAKARSPSPRPRRQSNGKATDTRGGNKVVDELKPKGASSSQSGSAAAAATAKRMAGSS KMRVIPSRYSLTPGASLGSSGAQERRRKQSLPGSSGDANQNEEIRAKVIEPSNDPLSPQTIS KVAEMLPKIRTMPPPDESPRDSGCAKRVAELVGKRSFFTAAAEDGRALDVEAPEAVAEA

# SEQ ID NO 5: Medicago trunculata seedyl encoding sequence

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# SEQ ID NO 6: Medicago trunculata seedyl protein

MNNTNNNNILLHSTQVQVWNNAAFDGEDFAMNSSSDSIKENLNPSAFNIVPSSNKRTIDDEI AEIESEIKRLTSKLELLRVEKAERKIASEKRVSGIGTGRIVAAKFMEPKKNVTPKRNGVVFK EETPKRNGVVSDTPKSRVNWRRGMSLGPMEIAGKVMAPPAMTITPATVNRRKSCFWKPQESC EVMPSGITPATVNRRKSCFLKPQESCEENRRKTICKPNLNLNSNSVNSAVGSIKRVKKKDEE IAQVQPKKLFEGEKSVKKSLKQGRIVASRYNSGGGGGDARKRSFSENNKGLGSEIRAKKRWE IPIEEVDVSGFVMLPKISTMRFVDESPRDSGAVKRVAELNGKRSYFCDEDEEERVMVEEEGG SVCQVLNFAEDDDDDDDYGEQG

SEQ ID NO 8: Saccharum sp. seedyl protein (partial N-term)
MEEDPLIPLVHVWNNAAFDHASSSAWHAHSPVPASARREAEGDKENHRPDPDPDVEAEIGHI
EAEILRLXSRLHHLRTSKQSEPSKRGEVAPAPAAKAKAAAAARLRTRGLSLGPLDVAAAGNP
NPLTTDNQQQQPRAAQGLKPIKQATAAAGKGVRLGPLRHGRR

SEQ ID NO 9: Zea Mays seedy1 encoding sequence (partial 3'end) ccacgcgtccggccgttcgagaggaaggccagcgttccaaggagcacgccgtccccgcc agaccgtggccatccagcaatgccaggcacccactggatgccaggcaaggcaccgcagcaag caqaqccaaqqcqaqqqqqqqqaqcataaqccccaqcaqqttcaqqaqqcaqtccacttcca gcggtcaatcacaccagcaatgtagccacgacgaagaggccggcggggagctccaaggtcag ggttgtcccgagccgctacagcatcccacctggctcctccctagcagctgtgacacaaggca accgatgcaagcagtctctcccaggatcggctactgagaccagagtaaatctcactgagccq ccgaacgacgagttgtctcctgaagaacttgccaaggttgcagagctgctcccaaggattaq gaccatgccgccttctgatgagagcccgcgtgactcgggatgtgccaagcgtgttgctgatt tggtcgggaagcgatccttcttcactgctgcaggggacgatggcaatctcgttacgccctac caggcacgggtggttgaacttgaatcacccgaggcagcagcagaagaagcagaagcttgaga agtttgtctttgatcaattccgaagtggcttgcatctgggcgtggcctcttttttgcagtgtg agacattgcttagtacttttgtgtttgccttgtgaaaagagtggaaggttcatctgctgat nccttgtt

SEQ ID NO 10: Zea Mays seedy1 protein (partial C-term)
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# SEQ ID NO 11: Arabidospis thaliana seedyl coding sequence

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# SEQ ID NO 12: Arabidospis thaliana seedy1 protein

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# SEQ ID NO 13: Sequence of the [PRO0090 - CDS0689 - terminator] expression cassette

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#### SEQ ID NO 14: rice prolamin RP6 promoter sequence

SEQ ID NO 15: Motif 1 CORE SEQUENCE WXNAXXD

SEQ ID NO 16: Motif 2 CORE SEQUENCE KENXXP

SEQ ID NO 17: Motif 3 (coiled coil) CORE SEQUENCE  $EX_{1-6}EXXRLXXXLXXLR$ 

SEQ ID NO 18: Motif 4 CORE SEQUENCE LPXIX<sub>1-10</sub>RDSGXXKRX<sub>1-6</sub>K

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# 105-Seedyl-EP.ST25.txt

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Ser	Leu	Lys 35	Arg	Ser	Trp	Ser	Pro 40	Leu	Lys	Pro	Leu	Ser 45	Val	Arg	Pro
Ser	Asp 50	Ser	Phe	Glu	Ser	Asp 55	Leu	Ser	Ser	Lys	Glu 60	Asn	Gln	Thr	Pro
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Pro	Leu	Asn	Pro	Asn 85	Gly	Ala	Leu	Glu	Asn 90	Ser	Arg	Leu	Lys	Pro 95	Asn
Lys	Pro	Asn	Ser 100	Lys	Gln	Ser	Leu	Asp 105	Glu	Met	Ala	Ala	Arg 110	Lys	Ser
Gly	Lys	Gly 115	Asn	Asp	Phe	Arg	Asp 120	Glu	Lys	Lys	Ile	Asp 125	Glu	Glu	Ile
Glu	Glu 130	Ile	Gln	Met	Glu	Ile 135	Ser	Arg	Leu	Ser	Ser 140	Arg	Leu	Glu	Ala
Leu 145	Arg	Ile	Glu	Lys	Ala 150	Glu	Lys	Thr	Val	Ala 155	Lys	Thr	Val	Glu	Lys 160
Arg	Gly	Arg	Val	Val 165	Ala	Ala	Lys	Phe	Met 170	Glu	Pro	Lys	Gln	Ser 175	Val
Ile	Lys	Ile	Glu 180	Glu	Arg	Ile	Ser	Met 185	Ser	Ala	Arg	Thr	Lys 190	Val	Glu
Gln	Arg	Arg 195	Gly	Leu	Ser	Leu	Gly 200	Pro	Ser	Glu	Ile	Phe 205	Thr	Gly	Thr
Arg	Arg 210	Arg	Ģly	Ļeu	Ser	Met 215	Gly	Pro	Ser	Asp	Ile 220	Leu	Ala	Gly	Thr
Thr 225	Lys	Ala	Arg	Gln	Leu 230	Gly	Lys	Gln	Glu	Met 235	Ile	Ile	Thr	Pro	Ile 240
Gln	Pro	Ile	Gln	Asn 245	Arg	Arg	Lys	Ser	Cys 250	Phe	Trp	Lys	Leu	Gln 255	Glu
Ile	Glu	Glu	Glu	Gly	Lys	Ser	Ser	Ser	Leu	Ser	Pro	Lys	Ser	Arg	Lys

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260

Thr Ala Ala Arg Thr Met Val Thr Thr Arg Gln Ala Val Thr Thr Ile 275 280 285

Ala Ser Lys Lys Asn Leu Lys Lys Asp Asp Gly Leu Leu Ser Ser Val 290 295 300

Gln Pro Lys Lys Leu Phe Lys Asp Leu Glu Lys Ser Ala Ala Ala Asn 305 310 315 320

Lys Lys Pro Gln Arg Pro Gly Arg Val Val Ala Ser Arg Tyr Asn Gln 325 330 335

Ser Thr Ile Gln Ser Ser Val Val Arg Lys Arg Ser Leu Pro Glu Asn 340 345 350

Asp Lys Asp Glu Ser Lys Arg Asn Asp Lys Lys Arg Ser Leu Ser Val 355 360 365

Gly Lys Thr Arg Val Ser Gln Thr Glu Ser Lys Asn Leu Gly Thr Glu 370 375 380

Ser Arg Val Lys Lys Arg Trp Glu Ile Pro Ser Glu Ile Val Val His 385 390 395 400

Gly Asn Thr Glu Ser Glu Lys Ser Pro Leu Ser Ile Ile Val Lys Pro 405 410 415

Asp Leu Leu Pro Arg Ile Arg Ile Ala Arg Cys Val Asn Glu Thr Leu 420 425 430

Arg Asp Ser Gly Pro Ala Lys Arg Met Ile Glu Leu Ile Gly Lys Lys 435 440 445

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<210> 3

<211> 1336

# 105-Seedy1-EP.ST25.txt

<212> DNA

<213> Oryza sativa

<220>

<221> misc\_feature

<223> seedyl coding sequence

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cgcgtctcca	agcagccgga	gcccaaccgc	gacgacgctc	cgatggggga	gatggtcgcg	300
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gcaccagcgg	cggcgcgagc	caagccgttc	accaccaage	tcaacgccat	tcgagaagaa	660
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gccactgcca	agaggatggc	ggggagctcc	aagatgaggg	tcatcccgag	ccgctacagc	960
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ccttccaatg	atccactctc	tcctcaaacg	atctccaagg	ttgctgaaat	gctcccaaag	1140
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gccgaattgg	tegggaageg	ctcgttcttc	acggctgcag	ccgaggacgg	gcgggcgctc	1260
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<210> 4

<211> 431

<212> PRT

<213> Oryza sativa

<220>

<221> MISC\_FEATURE

<223> seedyl protein

<400> 4

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Ala Phe Asp Asp Ser Ser Cys Ser Arg Ser Ala Trp Leu Pro Gln Ser 25

Pro Ala Val Ala Val Arg Lys Gly Asp Lys Glu Asn His Arg Pro 35 40 45

Glu Val Val Asp Val Ala Ala Gly Tyr Asp Val Glu Ala Glu Ile Gly 50 55

His Ile Glu Ala Glu Ile Leu Arg Leu Ser Ser Arg Leu His His Leu 65 70 80

Arg Val Ser Lys Gln Pro Glu Pro Asn Arg Asp Asp Ala Pro Met Gly 95

Glu Met Val Ala Lys Val Arg Pro Arg Pro Arg Gly Leu Ser Leu Gly 100 100 110

Pro Leu Asp Val Ile Ser Ile Val Asn Arg Glu Lys His Pro Leu Arg 115 120 125

Thr Lys Gln Pro Pro Ala Thr Arg Gly Arg Gly Leu Ser Leu Gly Pro 130 135

			105-Seed	y1-EP.ST	25.txt	
Met Glu Ile A 145	la Ala Ala 150	Asn Pro	Arg Val	Pro Ala 155	Ala Ala	Gln His 160
Gln Gln Gln G	ln Arg Ala 165	Gly Thr	Ala Arg 170	Ile Leu	Lys Pro	Ile Lys 175
Glu Pro Pro V	al Gln Arg 80	Arg Arg	Gly Val 185	Ser Leu	Gly Pro 190	Leu Glu
Ile His His G	ly Val Gly	Ser Lys 200	Ala Pro	Ala Ala	Ala Arg 205	Ala Lys
Pro Phe Thr T	hr Lys Leu	Asn Ala 215	Ile Arg	Glu Glu 220	Thr Arg	Pro Ser
Lys Gln Phe A	la Val Pro 230	Ala Lys	Pro Trp	Pro Ser 235	Ser Asn	Thr Arg 240
Gln Thr Leu A	sp Ser Arg 245	Gln Gly	Thr Ala 250	Ala Ser	Arg Ala	Lys Ala 255
Arg Ser Pro S	er Pro Arg 60	Pro Arg	Arg Gln 265	Ser Asn	Gly Lys 270	Ala Thr
Asp Thr Arg G 275	ly Gly Asn	Lys Val 280	Val Asp	Glu Leu	Lys Pro 285	Lys Gly
Ala Ser Ser S 290	er Gln Ser	Gly Ser 295	Ala Ala	Ala Ala 300	Ala Thr	Ala Lys
Arg Met Ala G 305	ly Ser Ser 310	Lys Met	Arg Val	Ile Pro 315	Ser Arg	Tyr Ser 320
Leu Thr Pro G	ly Ala Ser 325	Leu Gly	Ser Ser 330	Gly Ala	Gln Glu	Arg Arg 335
Arg Lys. Gln S	er Leu Pro 40	Gly Ser	Ser Gly 345	Asp Ala	Asn Gln 350	Asn Glu
Glu Ile Arg A 355	la Lys Val	Ile Glu 360		Asn Asp	Pro Leu 365	Ser Pro
Gln Thr Ile S 370	er Lys Val	Ala Glu 375	Met Leu	Pro Lys 380	Ile Arg	Thr Met

#### 105-Seedyl-EP.ST25.txt

Pro Pro Pro Asp Glu Ser Pro Arg Asp Ser Gly Cys Ala Lys Arg Val 385 390 395 400

Ala Glu Leu Val Gly Lys Arg Ser Phe Phe Thr Ala Ala Ala Glu Asp 405 410 415

Gly Arg Ala Leu Asp Val Glu Ala Pro Glu Ala Val Ala Glu Ala 420 425 430

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<211> 1860

<212> DNA

<213> Medicago trunculata

<220>

<221> misc feature

<223> seedy1 coding sequence

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# 105-Seedyl-EP.ST25.txt

. :-

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gcaggaaagt	tgtgaagtaa	tgccgtcggg	gattactccg	gcgacggtga	ataggaggaa	960
atcttgtttt	ttgaaacctc	aagaaagttg	tgaagaaaat	cgaagaaaaa	cgatttgcaa	1020
accgaatttg	aatttgaatt	caaattcagt	taattctgcg	gttggatcga	ttaagcgtgt	1080
gaagaagaaa	gatgaagaaa	ttgctcaggt	tcaaccgaag	aagctgtttg	aaggtgaaaa	1140
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tggtggtggt	gatgcgagga	aaagatcgtt	ttcggagaat	aataagggtt	tagggagtga	1260
aatcagggct	aagaagagat	gggagatacc	aattgaagaa	gtggatgtga	gtggttttgt	1320
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tgttaaaaga	gttgctgaat	tgaatggaaa	aagatcttac	ttttgtgatg	aagatgagga	1440
ggagagagtg	atggtggagg	aagaaggtgg	ttctgtttgt	caggttttga	attttgctga	1500
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ttgtttttgt	ggggttgtgt	ggaactggct	atgttctgct	tgattctttt	gcattttggt	1620
gtgaaactaa	agatgaggtg	aaaagtttat	gcttgttaaa	ttggattggt	ttatatgttt	1680
tgaaataata	acaacaagca	tgtgtcttgc	ttaataattg	tatattgttt	tgtttgtttt	1740
ataatgatat	ggatttaatt	tgtatacaca	atataatata	gtatgcattg	agagagtttt	1800
tcgttcagta	ttcattctga	ttttagtgtt	tatctcattc	tagaagattg	tattttgttg	1860

<210> 6

<211> 394

<212> PRT

<213> Medicago trunculata

<220>

<221> MISC\_FEATURE

<223> seedy1 protein

<400> 6

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#### 105-Seedy1-EP.ST25.txt

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- Ser Ser Ser Asp Ser Ile Lys Glu Asn Leu Asn Pro Ser Ala Phe Asn 35 40 45
- Ile Val Pro Ser Ser Asn Lys Arg Thr Ile Asp Asp Glu Ile Ala Glu 50 55 60
- Ile Glu Ser Glu Ile Lys Arg Leu Thr Ser Lys Leu Glu Leu Leu Arg 65 70 75 80
- Val Glu Lys Ala Glu Arg Lys Ile Ala Ser Glu Lys Arg Val Ser Gly 85 90 95
- Ile Gly Thr Gly Arg Ile Val Ala Ala Lys Phe Met Glu Pro Lys Lys
  100 105 110
- Asn Val Thr Pro Lys Arg Asn Gly Val Val Phe Lys Glu Glu Thr Pro 115 120 125
- Lys Arg Asn Gly Val Val Ser Asp Thr Pro Lys Ser Arg Val Asn Trp 130 135 140
- Arg Arg Gly Met Ser Leu Gly Pro Met Glu Ile Ala Gly Lys Val Met 145 150 155 160
- Ala Pro Pro Ala Met Thr Ile Thr Pro Ala Thr Val Asn Arg Arg Lys
  165 170 175
- Ser Cys Phe Trp Lys Pro Gln Glu Ser Cys Glu Val Met Pro Ser Gly 180 185 190
- Ile Thr Pro Ala Thr Val Asn Arg Arg Lys Ser Cys Phe Leu Lys Pro 195 200 205
- Gln Glu Ser Cys Glu Glu Asn Arg Arg Lys Thr Ile Cys Lys Pro Asn 210 215 220
- Leu Asn Leu Asn Ser Asn Ser Val Asn Ser Ala Val Gly Ser Ile Lys 225 230 235 240
- Arg Val Lys Lys Asp Glu Glu Ile Ala Gln Val Gln Pro Lys Lys 245 250 255

Leu Phe Glu Gly Glu Lys Ser Val Lys Lys Ser Leu Lys Gln Gly Arg 260 265 270

Ile Val Ala Ser Arg Tyr Asn Ser Gly Gly Gly Gly Asp Ala Arg
275 280 285

Lys Arg Ser Phe Ser Glu Asn Asn Lys Gly Leu Gly Ser Glu Ile Arg 290 295 300

Ala Lys Lys Arg Trp Glu Ile Pro Ile Glu Glu Val Asp Val Ser Gly 305 310 315 320

Phe Val Met Leu Pro Lys Ile Ser Thr Met Arg Phe Val Asp Glu Ser 325 330 335

Pro Arg Asp Ser Gly Ala Val Lys Arg Val Ala Glu Leu Asn Gly Lys 340 345 350

Arg Ser Tyr Phe Cys Asp Glu Asp Glu Glu Glu Arg Val Met Val Glu 355 360 365

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Asp Asp Asp Asp Tyr Gly Glu Gln Gly 385

<210> 7

<211> 674

<212> DNA

<213> Saccharum sp.

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<221> misc\_feature

<223> seedyl coding sequence (partial 5' end)

<220>

<221> misc\_feature

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<223> n can be a, c, g or t

<220>
<221> misc\_feature
<222> (372)..(372)
<223> n can be a, c, g or t

<220>
<221> misc\_feature
<222> (674)..(674)
<223> n can be a, c, g or t

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<210> 8

<211> 166

<212> PRT

<213> Saccharum sp.

<220>

<221> MISC\_FEATURE

<223> seedyl protein

<220>

<221> MISC FEATURE

<223> seedyl protein (partial N term)

<220>

<221> MISC\_FEATURE

<222> (70)..(70)

<223> Xaa can be any amino acid

<400> 8

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Ala Phe Asp His Ala Ser Ser Ser Ala Trp His Ala His Ser Pro Val 20 25 30

Pro Ala Ser Ala Arg Arg Glu Ala Glu Gly Asp Lys Glu Asn His Arg 35 40 45

Pro Asp Pro Asp Pro Asp Val Glu Ala Glu Ile Gly His Ile Glu Ala 50 55 60

Glu Ile Leu Arg Leu Xaa Ser Arg Leu His His Leu Arg Thr Ser Lys 65 70 75 80

Gln Ser Glu Pro Ser Lys Arg Gly Glu Val Ala Pro Ala Pro Ala Ala 85 90 95

Lys Ala Lys Ala Ala Ala Ala Ala Arg Leu Arg Thr Arg Gly Leu Ser 100 105 110

Leu Gly Pro Leu Asp Val Ala Ala Ala Gly Asn Pro Asn Pro Leu Thr 125 Thr Asp Asn Gln Gln Gln Pro Arg Ala Ala Gln Gly Leu Lys Pro 135 Ile Lys Gln Ala Thr Ala Ala Gly Lys Gly Val Arg Leu Gly Pro Leu Arg His Gly Arg Arg <210> 9 <211> 876 <212> DNA <213> Zea Mays <220> <221> misc feature seedyl coding sequence (partila 3' end) <223> <220> <221> misc\_feature <222> (869)..(869)  $\langle 223 \rangle$  n = a, c, g or t <400> ccacgcgtcc ggccgttcga gaggaggaag gccagcgttc caaggagcac gccgtccccg ccagaccgtg gccatccagc aatgccaggc acccactgga tgccaggcaa ggcaccgcag

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ccagaccgtg gccatccagc aatgccaggc acccactgga tgccaggcaa ggcaccgcag 120
caagcagagc caaggcgagg agcgggagca taagccccag caggttcagg aggcagtcca 180
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ggagcgaagc ggtcaatcac accagcaatg tagccacgac gaagaggccg gcggggagct 300
ccaaggtcag ggttgtcccg agccgctaca gcatcccacc tggctcctcc ctagcagctg 360
tgacacaagg caaccgatgc aagcagtctc tcccaggatc ggctactgag accagagtaa 420
Page 14

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ccaagcgtgt tgctga	tttg gtcgggaagc q	gatccttctt	cactgctgca	ggggacgatg	600
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ggcgtggcct cttttt	gcag tgtgtgctac 1	tacatagtct	actgttacat	tcatatcata	780
tcacatttcc tatttt	ttcc cccttgagac a	attgcttagt	acttttgtgt	tgccttgtga	840
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<210> 10

<211> 224

<212> PRT

<213> Zea Mays

<220>

<221> MISC\_FEATURE

<223> seedyl protein (partial C term)

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1 10 15

Ala Val Pro Ala Arg Pro Trp Pro Ser Ser Asn Ala Arg His Pro Leu 20 25 30

Asp Ala Arg Gln Gly Thr Ala Ala Ser Arg Ala Lys Ala Arg Ser Gly 35 40 45

Ser Ile Ser Pro Ser Arg Phe Arg Gln Ser Thr Ser Lys Ala Ala 50 55 60

Glu Thr Arg Ala Gly Asn Ala Lys Pro Thr Glu Ala Thr Arg Gly Gly 65 70 75 80

Ser Glu Ala Val Asn His Thr Ser Asn Val Ala Thr Thr Lys Arg Pro Page 15

85 ·	90	95

Ala Gly Ser Ser Lys Val Arg Val Val Pro Ser Arg Tyr Ser Ile Pro 100 105 110

Pro Gly Ser Ser Leu Ala Ala Val Thr Gln Gly Asn Arg Cys Lys Gln
115 120 125

Ser Leu Pro Gly Ser Ala Thr Glu Thr Arg Val Asn Leu Thr Glu Pro 130 135 140

Pro Asn Asp Glu Leu Ser Pro Glu Glu Leu Ala Lys Val Ala Glu Leu 145 150 155 160

Leu Pro Arg Ile Arg Thr Met Pro Pro Ser Asp Glu Ser Pro Arg Asp 165 170 175

Ser Gly Cys Ala Lys Arg Val Ala Asp Leu Val Gly Lys Arg Ser Phe 180 185 190

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<210> 11

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<212> DNA

<213> Arabidospis thaliana

<220>

<221> misc\_feature

<223> seedy1 coding sequence

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tctcacctca acgaatcatt cgattccgat tgtagcaagg agaatcagtt tccgatttcg 180

gtttcctctt	cgctccaatc	ctcagtctcg	atcaccgaag	ctccgtcagc	aaaatccaag	240
accgtgaaga	ccaaatccgc	cgcagatcgg	agtaaaaagc	gagatatcga	tgcagagatc	300
gaagaagtag	agaaggagat	cggacgatta	tcgacgaaat	tggagtcgct	ccgattagag	360
aaggcggagc	aaaccgcaag	aagcattgct	atacgtggaa	gaatcgttcc	ggcgaagttc	420
atggaatcat	ctcagaaaca	agtgaaattc	gacgattcgt	gttttacagg	atcgaaatca	480
agagccactc	gtagaggcgt	tagtcttgga	ccagcggaga	tattcaattc	cgcgaagaaa	540
tctgaaactg	tgactcctct	tcaatcagct	cagaatcgac	gcaagtcttg	tttctttaag	600
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gtggggtcaa	agagagctgt	gaagaaagaa	gaaggagttc	tcttaacaat	ccagcctaag	780
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gcttctgatg	aaagtaacaa	gagtgaaggg	agagtgaaga	agagatggga	gattccaagt	1020
gaagttgatc	tgtatagcag	tggtgagaac	ggtgacgagt	ctcctatagt	taaggagcta	1080
cctaagatca	gaacgcttcg	tcgtgtggga	gggagccctc	gtgattcagg	tgctgctaag	1140
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<210> 12

<211> 402

<212> PRT

<213> Arabidospis thaliana

<220> ... ... ... ...

<221> MISC\_FEATURE

<223> seedy1 protein

<400> 12

Met Thr Ser Ile Glu Ala Thr Glu Thr Leu Asn Ala Pro Pro Lys Leu Page 17

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Val Thr Thr Arg Gly Lys Gly Arg Thr Ser Leu Ser Leu Ser Pro Arg

Ser Arg Lys Ala Lys Met Thr Ala Ala Gln Lys Gln Ala Ala Thr Thr

230

235

240

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